

Welcome to STN International! Enter x:x

LOGINID:sssptau125rxt

PASSWORD:

TERMINAL (ENTER 1, 2, 3, OR ?):2

* * * * * * * * * * Welcome to STN International * * * * * * * * *

NEWS 1 Web Page URLs for STN Seminar Schedule - N. America
NEWS 2 "Ask CAS" for self-help around the clock
NEWS 3 Feb 24 PCTGEN now available on STN
NEWS 4 Feb 24 TEMA now available on STN
NEWS 5 Feb 26 NTIS now allows simultaneous left and right truncation
NEWS 6 Feb 26 PCTFULL now contains images
NEWS 7 Mar 04 SDI PACKAGE for monthly delivery of multifile SDI results
NEWS 8 Mar 24 PATDPAFULL now available on STN
NEWS 9 Mar 24 Additional information for trade-named substances without
structures available in REGISTRY
NEWS 10 Apr 11 Display formats in DGENE enhanced
NEWS 11 Apr 14 MEDLINE Reload
NEWS 12 Apr 17 Polymer searching in REGISTRY enhanced
NEWS 13 Jun 13 Indexing from 1947 to 1956 added to records in CA/CAPLUS
NEWS 14 Apr 21 New current-awareness alert (SDI) frequency in
WPIDS/WPINDEX/WPIX
NEWS 15 Apr 28 RDISCLOSURE now available on STN
NEWS 16 May 05 Pharmacokinetic information and systematic chemical names
added to PHAR
NEWS 17 May 15 MEDLINE file segment of TOXCENTER reloaded
NEWS 18 May 15 Supporter information for ENCOMPPAT and ENCOMPLIT updated
NEWS 19 May 19 Simultaneous left and right truncation added to WSCA
NEWS 20 May 19 RAPRA enhanced with new search field, simultaneous left and
right truncation
NEWS 21 Jun 06 Simultaneous left and right truncation added to CBNB
NEWS 22 Jun 06 PASCAL enhanced with additional data
NEWS 23 Jun 20 2003 edition of the FSTA Thesaurus is now available
NEWS 24 Jun 25 HSDB has been reloaded

NEWS EXPRESS April 4 CURRENT WINDOWS VERSION IS V6.01a, CURRENT
MACINTOSH VERSION IS V6.0b(ENG) AND V6.0Jb(JP),
AND CURRENT DISCOVER FILE IS DATED 01 APRIL 2003
NEWS HOURS STN Operating Hours Plus Help Desk Availability
NEWS INTER General Internet Information
NEWS LOGIN Welcome Banner and News Items
NEWS PHONE Direct Dial and Telecommunication Network Access to STN
NEWS WWW CAS World Wide Web Site (general information)

Enter NEWS followed by the item number or name to see news on that
specific topic.

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of commercial gateways or other similar uses is prohibited and may
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* * * * * * * * * * STN Columbus * * * * * * * * * * * * * * *

FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003

| | | | |
|----------------------|--|------------|---------|
| => file reg | | | |
| COST IN U.S. DOLLARS | | SINCE FILE | TOTAL |
| | | ENTRY | SESSION |
| FULL ESTIMATED COST | | 0.21 | 0.21 |

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003
USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT.
PLEASE SEE "HELP USAGETERMS" FOR DETAILS.
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Property values tagged with IC are from the ZIC/VINITI data file
provided by InfoChem.

STRUCTURE FILE UPDATES: 14 JUL 2003 HIGHEST RN 548428-18-8
DICTIONARY FILE UPDATES: 14 JUL 2003 HIGHEST RN 548428-18-8

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when
conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP
PROPERTIES for more information. See STNote 27, Searching Properties
in the CAS Registry File, for complete details:
<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

=> scedrol
SCEDROL IS NOT A RECOGNIZED COMMAND
The previous command name entered was not recognized by the system.
For a list of commands available to you in the current file, enter
"HELP COMMANDS" at an arrow prompt (=>).

=> s cedrol
L1 43 CEDROL

=> s patchouli
L2 33 PATCHOULI

=> s santalol
L3 36 SANTALOL

=> s l3 36
MISSING OPERATOR

=> d l3 36

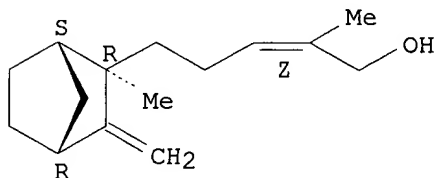
L3 ANSWER 36 OF 36 REGISTRY COPYRIGHT 2003 ACS
RN 77-42-9 REGISTRY
CN 2-Penten-1-ol, 2-methyl-5-[(1S,2R,4R)-2-methyl-3-
methylenebicyclo[2.2.1]hept-2-yl]-, (2Z)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:

CN .beta.-Santalol (6CI)
CN 2-Penten-1-ol, 2-methyl-5-(2-methyl-3-methylene-2-norbornyl)- (7CI, 8CI)
CN 2-Penten-1-ol, 2-methyl-5-(2-methyl-3-methylenebicyclo[2.2.1]hept-2-yl)-,
[1S-[1.alpha.,2.alpha.(Z),4.alpha.]]-

OTHER NAMES:

CN (-)-(Z)-.beta.-Santalol
CN (-).beta.-Santalol
CN cis-.beta.-Santalol
CN Santalol b
FS STEREOSEARCH
DR 37172-31-9
MF C15 H24 O
LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA,
CAOLD, CAPLUS, CASREACT, CHEMLIST, CIN, HODOC*, HSDB*, IFICDB, IFIPAT,
IFIUDB, MRCK*, NAPRALERT, SPECINFO, TOXCENTER, USPATFULL
(*File contains numerically searchable property data)
Other Sources: DSL**, EINECS**, TSCA**
(**Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.
Double bond geometry as shown.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

129 REFERENCES IN FILE CA (1957 TO DATE)
4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
129 REFERENCES IN FILE CAPLUS (1957 TO DATE)
9 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

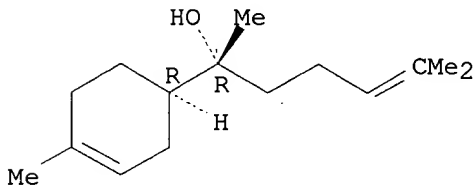
=> s bisabolol
L4 31 BISABOLOL

=> d 14 31

L4 ANSWER 31 OF 31 REGISTRY COPYRIGHT 2003 ACS
RN 515-69-5 REGISTRY
CN 3-Cyclohexene-1-methanol, .alpha.,4-dimethyl-.alpha.- (4-methyl-3-pentenyl)-
, (.alpha.R,1R)-rel- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN 3-Cyclohexene-1-methanol, .alpha.,4-dimethyl-.alpha.- (4-methyl-3-pentenyl)-
, (R*,R*)-
CN 5-Hepten-2-ol, 6-methyl-2-(4-methyl-3-cyclohexen-1-yl)- (6CI, 7CI, 8CI)
OTHER NAMES:
CN (+-).alpha.-Bisabolol
CN .alpha.-Bisabolol
CN Bisabolol
CN Camilol
CN dl-.alpha.-Bisabolol
CN Dragosantol
CN Hydagen B
FS STEREOSEARCH
DR 63601-23-0, 25428-43-7, 21090-60-8, 67375-41-1
MF C15 H26 O
CI COM

LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS,
BIOTECHNO, CA, CABA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CHEMCATS,
CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, EMBASE, IPA, MEDLINE, MRCK*, PIRA,
PROMT, RTECS*, SPECINFO, TOXCENTER, USPAT2, USPATFULL
(*File contains numerically searchable property data)
Other Sources: DSL**, EINECS**, TSCA**
(**Enter CHEMLIST File for up-to-date regulatory information)

Relative stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

857 REFERENCES IN FILE CA (1957 TO DATE)
11 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
863 REFERENCES IN FILE CAPLUS (1957 TO DATE)
10 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> e bisabolol
E1 9 BISABOLENOL/BI
E2 1 BISABOLIDE/BI
E3 31 --> BISABOLOL/BI
E4 1 BISABOLOLONE/BI
E5 2 BISABOLON/BI
E6 6 BISABOLONE/BI
E7 2 BISABOLONOXIDE/BI
E8 1 BISABOLOXIDE/BI
E9 1 BISABOLYL/BI
E10 2 BISABON/BI
E11 1 BISABONE/BI
E12 2 BISABONOL/BI

=> s vetiverol
L5 2 VETIVEROL

=> d 15 1 2

L5 ANSWER 1 OF 2 REGISTRY COPYRIGHT 2003 ACS
RN 68129-81-7 REGISTRY
CN **Vetiverol (9CI)** (CA INDEX NAME)
OTHER NAMES:
CN Lignolia
CN Vetivenol
CN Vetivol
MF Unspecified
CI COM, MAN
LC STN Files: BIOBUSINESS, BIOSIS, CA, CAPLUS, CHEMCATS, CHEMLIST, CIN,
CSCHEM, DDFU, DRUGU, NAPRALERT, RTECS*, TOXCENTER, USPAT2, USPATFULL
(*File contains numerically searchable property data)
Other Sources: DSL**, EINECS**

(**Enter CHEMLIST File for up-to-date regulatory information)

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

53 REFERENCES IN FILE CA (1957 TO DATE)

4 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

53 REFERENCES IN FILE CAPLUS (1957 TO DATE)

L5 ANSWER 2 OF 2 REGISTRY COPYRIGHT 2003 ACS

RN 62563-80-8 REGISTRY

CN **Vetiverol, acetate (6CI, 9CI)** (CA INDEX NAME)

OTHER NAMES:

CN Vetiveryl acetate

MF C2 H4 O2 . x Unspecified

LC STN Files: BIOSIS, CA, CAOLD, CAPLUS, CHEMCATS, CHEMLIST, CIN, CSCHEM, RTECS*, TOXCENTER, USPATFULL

(*File contains numerically searchable property data)

Other Sources: DSL**, EINECS**

(**Enter CHEMLIST File for up-to-date regulatory information)

CM 1

CRN 68129-81-7

CMF Unspecified

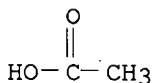
CCI MAN

*** STRUCTURE DIAGRAM IS NOT AVAILABLE ***

CM 2

CRN 64-19-7

CMF C2 H4 O2



38 REFERENCES IN FILE CA (1957 TO DATE)

38 REFERENCES IN FILE CAPLUS (1957 TO DATE)

3 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> s sclareol

L6 30 SCLAREOL

=> d 16 30

L6 ANSWER 30 OF 30 REGISTRY COPYRIGHT 2003 ACS

RN 515-03-7 REGISTRY

CN 1-Naphthalenopropanol, .alpha.-ethenyldecahydro-2-hydroxy-.alpha.,2,5,5,8a-pentamethyl-, (.alpha.R,1R,2R,4aS,8aS)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1-Naphthalenopropanol, .alpha.-ethenyldecahydro-2-hydroxy-.alpha.,2,5,5,8a-pentamethyl-, [1R-[1.alpha.(R*),2.beta.,4a.beta.,8a.alpha.]]-

CN Labd-14-ene-8,13-diol, (13R)- (8CI)

CN **Sclareol (6CI)**

OTHER NAMES:

CN **(-)-Sclareol**

FS STEREOSEARCH

DR 17904-64-2

MF C20 H36 O2

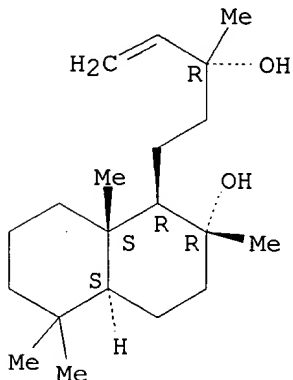
CI COM

LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS,
BIOTECHNO, CA, CANCERLIT, CAOLD, CAPLUS, CASREACT, CBNB, CHEMCATS,
CHEMINFORMRX, CHEMLIST, CIN, CSCHEM, DDFU, DRUGU, EMBASE, MEDLINE,
NAPRALERT, NIOSHTIC, PROMT, RTECS*, SPECINFO, TOXCENTER, USPATFULL
(*File contains numerically searchable property data)

Other Sources: DSL**, EINECS**, TSCA**

(**Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry. Rotation (-).



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

261 REFERENCES IN FILE CA (1957 TO DATE)

1 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA

261 REFERENCES IN FILE CAPLUS (1957 TO DATE)

19 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> d 16 29

L6 ANSWER 29 OF 30 REGISTRY COPYRIGHT 2003 ACS

RN 564-20-5 REGISTRY

CN Naphtho[2,1-b]furan-2(1H)-one, decahydro-3a,6,6,9a-tetramethyl-,
(3aR,5aS,9aS,9bR)- (9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN Naphtho[2,1-b]furan-2(1H)-one, 3a,4,5,5a.alpha.,6,7,8,9,9a,9b.alpha.-
decahydro-3a.beta.,6,6,9a.beta.-tetramethyl- (8CI)

CN Naphtho[2,1-b]furan-2(1H)-one, decahydro-3a,6,6,9a-tetramethyl-,
[3aR-(3a.alpha.,5a.beta.,9a.alpha.,9b.beta.)]-

CN Norambreinolide (6CI, 7CI)

OTHER NAMES:

CN (+)-Norambreinolide

CN (+)-Sclareolide

CN Norambreinolid

CN Sclareolide

FS STEREOSEARCH

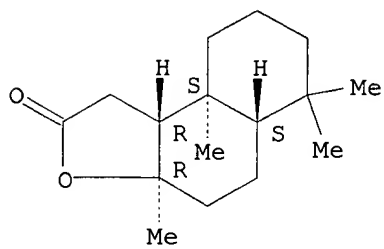
MF C16 H26 O2

LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA,
CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CIN, CSCHEM,
NAPRALERT, SPECINFO, TOXCENTER, USPATFULL
(*File contains numerically searchable property data)

Other Sources: DSL**, EINECS**, TSCA**

(**Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry. Rotation (+).



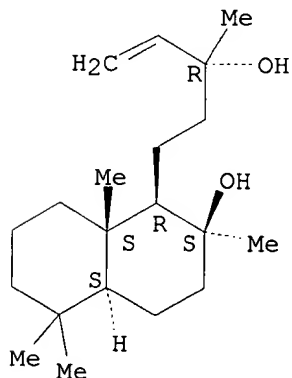
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

140 REFERENCES IN FILE CA (1957 TO DATE)
140 REFERENCES IN FILE CAPLUS (1957 TO DATE)
5 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> d 16 28

L6 ANSWER 28 OF 30 REGISTRY COPYRIGHT 2003 ACS
RN 1232-00-4 REGISTRY
CN 1-Naphthalenopropanol, .alpha.-ethenyldecahydro-2-hydroxy-.alpha.,2,5,5,8a-pentamethyl-, [1R-[1.alpha.(R*),2.alpha.,4a.beta.,8a.alpha.]]- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN Labd-14-ene-8.beta.,13-diol (7CI, 8CI)
OTHER NAMES:
CN **8-Episclareol**
CN 8.beta.-Labd-14-ene-8,13-diol
FS STEREOSEARCH
MF C20 H36 O2
LC STN Files: BEILSTEIN*, CA, CAOLD, CAPLUS, CASREACT, CHEMINFORMRX
(*File contains numerically searchable property data)

Absolute stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

4 REFERENCES IN FILE CA (1957 TO DATE)
4 REFERENCES IN FILE CAPLUS (1957 TO DATE)
2 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

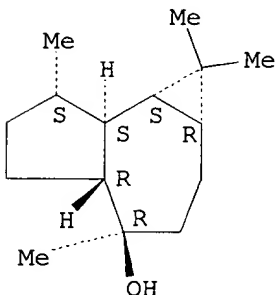
=> s globuol
L7 0 GLOBUOL

=> s globulol
L8 9 GLOBULOL

=> d 18 7 8 9

L8 ANSWER 7 OF 9 REGISTRY COPYRIGHT 2003 ACS
RN 55659-76-2 REGISTRY
CN 1H-Cycloprop[e]azulen-4-ol, decahydro-1,1,4,7-tetramethyl-,
(1aR,4R,4aR,7S,7aS,7bS)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN 1H-Cycloprop[e]azulen-4-ol, decahydro-1,1,4,7-tetramethyl-,
[1aR-(1a.alpha.,4.alpha.,4a.alpha.,7.beta.,7a.beta.,7b.alpha.)]-
OTHER NAMES:
CN (-)-4-Epiglobulol
CN 4-Epiglobulol, (-)-
FS STEREOSEARCH
MF C15 H26 O
LC STN Files: BEILSTEIN*, CA, CAPLUS, CHEMINFORMRX
(*File contains numerically searchable property data)

Absolute stereochemistry.



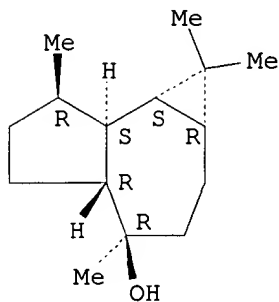
PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

4 REFERENCES IN FILE CA (1957 TO DATE)
4 REFERENCES IN FILE CAPLUS (1957 TO DATE)

L8 ANSWER 8 OF 9 REGISTRY COPYRIGHT 2003 ACS
RN 51371-47-2 REGISTRY
CN 1H-Cycloprop[e]azulen-4-ol, decahydro-1,1,4,7-tetramethyl-,
(1a.alpha.,4.alpha.,4a.alpha.,7.alpha.,7a.beta.,7b.alpha.)- (9CI) (CA
INDEX NAME)
OTHER CA INDEX NAMES:
CN 1H-Cycloprop[e]azulen-4-ol, decahydro-1,1,4,7-tetramethyl-,
(1a.alpha.,4.alpha.,4a.alpha.,7.alpha.,7a.beta.,7b.alpha.)-(-.+.-)-
OTHER NAMES:
CN (-)-Globulol
FS STEREOSEARCH
MF C15 H26 O

LC STN Files: BEILSTEIN*, CA, CAPLUS, CHEMCATS, CHEMINFORMRX
(*File contains numerically searchable property data)

Relative stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1957 TO DATE)
3 REFERENCES IN FILE CAPLUS (1957 TO DATE)

L8 ANSWER 9 OF 9 REGISTRY COPYRIGHT 2003 ACS

RN 489-41-8 REGISTRY

CN 1H-Cycloprop[e]azulen-4-ol, decahydro-1,1,4,7-tetramethyl-,
(1aR,4R,4aR,7R,7aS,7bS)- (8CI, 9CI) (CA INDEX NAME)

OTHER CA INDEX NAMES:

CN 1H-Cycloprop[e]azulen-4-ol, decahydro-1,1,4,7-tetramethyl-,
[1aR-(1a.alpha.,4.alpha.,4a.alpha.,7.alpha.,7a.beta.,7b.alpha.)]-

CN Globulol (6CI, 7CI)

OTHER NAMES:

CN (-)-Globulol

FS STEREOSEARCH

MF C15 H26 O

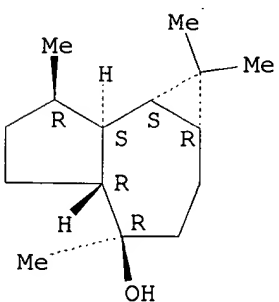
LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA,
CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CSCHEM,
NAPRALERT, SPECINFO, TOXCENTER, USPAT2, USPATFULL

(*File contains numerically searchable property data)

Other Sources: EINECS**

(**Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

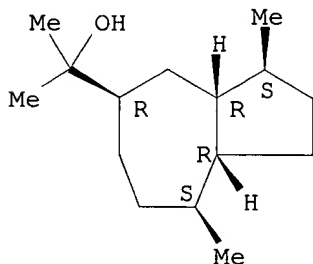
592 REFERENCES IN FILE CA (1957 TO DATE)
594 REFERENCES IN FILE CAPLUS (1957 TO DATE)
14 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

=> s guaiol
L9 11 GUAIOL

=> d 19 8 9 10 11

L9 ANSWER 8 OF 11 REGISTRY COPYRIGHT 2003 ACS
RN 3526-76-9 REGISTRY
CN 5-Azulenemethanol, decahydro-.alpha.,.alpha.,3,8-tetramethyl-,
[3S-(3.alpha.,3a.alpha.,5.alpha.,8.alpha.,8a.alpha.)]- (9CI) (CA INDEX
NAME)
OTHER CA INDEX NAMES:
CN 1.beta.,5.beta.-Guaien-11-ol (8CI)
OTHER NAMES:
CN 1.**beta.**,5.**beta.**-Dihydroguaiol
FS STEREOSEARCH
MF C15 H28 O
LC STN Files: BEILSTEIN*, CA, CAOLD, CAPLUS
(*File contains numerically searchable property data)

Absolute stereochemistry.

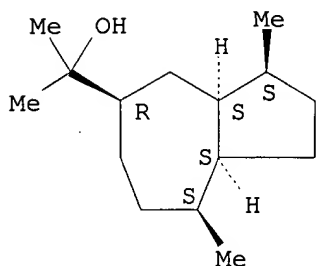


PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

3 REFERENCES IN FILE CA (1957 TO DATE)
3 REFERENCES IN FILE CAPLUS (1957 TO DATE)
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L9 ANSWER 9 OF 11 REGISTRY COPYRIGHT 2003 ACS
RN 3526-75-8 REGISTRY
CN 5-Azulenemethanol, decahydro-.alpha.,.alpha.,3,8-tetramethyl-,
(3S,3aS,5R,8S,8aS)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN 5-Azulenemethanol, decahydro-.alpha.,.alpha.,3,8-tetramethyl-,
[3S-(3.alpha.,3a.beta.,5.alpha.,8.alpha.,8a.beta.)]-
CN Guaien-11-ol (8CI)
OTHER NAMES:
CN 1.**alpha.**,5.**alpha.**-Dihydroguaiol
CN Galbanol
FS STEREOSEARCH
MF C15 H28 O
LC STN Files: BEILSTEIN*, CA, CAOLD, CAPLUS, CHEMLIST
(*File contains numerically searchable property data)
Other Sources: DSL**, EINECS**, TSCA**
(**Enter CHEMLIST File for up-to-date regulatory information)

Absolute stereochemistry.

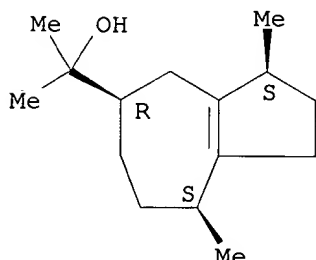


PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

6 REFERENCES IN FILE CA (1957 TO DATE)
6 REFERENCES IN FILE CAPLUS (1957 TO DATE)
1 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L9 ANSWER 10 OF 11 REGISTRY COPYRIGHT 2003 ACS
RN 489-86-1 REGISTRY
CN 5-Azulenemethanol, 1,2,3,4,5,6,7,8-octahydro-.alpha.,.alpha.,3,8-tetramethyl-, (3S,5R,8S)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN 5-Azulenemethanol, 1,2,3,4,5,6,7,8-octahydro-.alpha.,.alpha.,3,8-tetramethyl-, [3S-(3.alpha.,5.alpha.,8.alpha.)]-
CN 5-Azulenemethanol, 1,2,3,4,5.beta.,6,7,8-octahydro-.alpha.,.alpha.,3.alpha.,8.alpha.-tetramethyl- (7CI)
CN Guai-1(5)-en-11-ol (8CI)
CN **Guaiol (6CI)**
OTHER NAMES:
CN **(-) -Guaiol**
CN 3,8-Dimethyl-5-.alpha.-hydroxyisopropyl-.DELTA.9-octahydroazulene
CN Champaca camphor
CN Champacol
CN Guaiac alcohol
FS STEREOSEARCH
MF C15 H26 O
LC STN Files: AGRICOLA, ANABSTR, BEILSTEIN*, BIOBUSINESS, BIOSIS, CA, CAOLD, CAPLUS, CASREACT, CHEMCATS, CHEMINFORMRX, CHEMLIST, CSCHEM, DETHERM*, EMBASE, HODOC*, IPA, MRCK*, NAPRALERT, NIOSHTIC, SPECINFO, TOXCENTER, USPAT2, USPATFULL
(*File contains numerically searchable property data)
Other Sources: DSL**, EINECS**, TSCA**
(**Enter CHEMLIST File for up-to-date regulatory information)

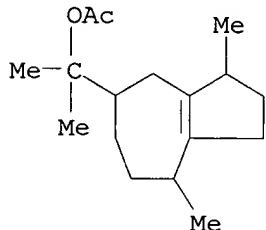
Absolute stereochemistry.



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

391 REFERENCES IN FILE CA (1957 TO DATE)
5 REFERENCES TO NON-SPECIFIC DERIVATIVES IN FILE CA
400 REFERENCES IN FILE CAPLUS (1957 TO DATE)
11 REFERENCES IN FILE CAOLD (PRIOR TO 1967)

L9 ANSWER 11 OF 11 REGISTRY COPYRIGHT 2003 ACS
RN 134-28-1 REGISTRY
CN 5-Azulenemethanol, 1,2,3,4,5,6,7,8-octahydro-.alpha.,.alpha.,3,8-tetramethyl-, acetate, (3S,5R,8S)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN 5-Azulenemethanol, 1,2,3,4,5,6,7,8-octahydro-.alpha.,.alpha.,3,8-tetramethyl-, acetate, [3S-(3.alpha.,5.alpha.,8.alpha.)]-
CN 5-Azulenemethanol, 1,2,3,4,5.alpha.,6,7,8-octahydro-.alpha.,.alpha.,3.beta.,8.beta.-tetramethyl-, acetate (8CI)
OTHER NAMES:
CN Guaiac acetate
CN **Guaiol acetate**
CN Guaiyl acetate
DR 17431-48-0
MF C17 H28 O2
LC STN Files: CA, CAPLUS, CHEMCATS, CHEMLIST, CSCHEM, SPECINFO, TOXCENTER
Other Sources: DSL**, EINECS**, TSCA**
(**Enter CHEMLIST File for up-to-date regulatory information)



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

21 REFERENCES IN FILE CA (1957 TO DATE)
21 REFERENCES IN FILE CAPLUS (1957 TO DATE)

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

| | |
|----|----------------|
| L1 | 43 S CEDROL |
| L2 | 33 S PATCHOULI |
| L3 | 36 S SANTALOL |
| L4 | 31 S BISABOLOL |
| | E BISABOLOL |
| L5 | 2 S VETIVEROL |
| L6 | 30 S SCLAREOL |
| L7 | 0 S GLOBUOL |
| L8 | 9 S GLOBULOL |
| L9 | 11 S GUIOL |

=> file caplus

| COST IN U.S. DOLLARS | SINCE FILE ENTRY | TOTAL SESSION |
|----------------------|------------------|---------------|
| FULL ESTIMATED COST | 65.50 | 65.71 |

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003
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FILE COVERS 1907 - 15 Jul 2003 VOL 139 ISS 3
 FILE LAST UPDATED: 14 Jul 2003 (20030714/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

```
=> e sleep
E1          1      SLEENOPHENE/BI
E2          1      SLEEO/BI
E3      15616 --> SLEEP/BI
E4          1      SLEEP10/BI
E5          1      SLEEP2/BI
E6          1      SLEEP9ING/BI
E7          1      SLEEPAGE/BI
E8          1      SLEEPDEPRIVED/BI
E9          1      SLEEPED/BI
E10     161      SLEEPER/BI
E11     199      SLEEPERS/BI
E12          4      SLEEPIER/BI
```

```
=> s e3
L10      15616 SLEEP/BI
```

```
=> e sedative
E1          1      SEDATIVA/BI
E2          1      SEDATIVAE/BI
E3      9549 --> SEDATIVE/BI
E4          1      SEDATIVEACTION/BI
E5          1      SEDATIVEFOR/BI
E6          4      SEDATIVEHYPNOTIC/BI
E7          3      SEDATIVELIKE/BI
E8          3      SEDATIVELY/BI
E9      7525      SEDATIVES/BI
E10     1       SEDATIVESC/BI
E11     1       SEDATIVUM/BI
E12     1       SEDATIVUS/BI
```

```
=> s e3-e9
9549 SEDATIVE/BI
1 SEDATIVEACTION/BI
1 SEDATIVEFOR/BI
```

4 SEDATIVEHYPNOTIC/BI
3 SEDATIVELIKE/BI
3 SEDATIVELY/BI
7525 SEDATIVES/BI
L11 12852 (SEDATIVE/BI OR SEDATIVEACTION/BI OR SEDATIVEFOR/BI OR SEDATIVEHYPNOTIC/BI OR SEDATIVELIKE/BI OR SEDATIVELY/BI OR SEDATIVES/BI)

=> e relaxation

E1 1 RELAXATIOIN/BI
E2 2 RELAXATIOIM/BI
E3 216874 --> RELAXATION/BI
E4 2 RELAXATION1/BI
E5 1 RELAXATION50/BI
E6 1 RELAXATIONA/BI
E7 1 RELAXATIONAAOF/BI
E8 2096 RELAXATIONAL/BI
E9 7 RELAXATIONALLY/BI
E10 1 RELAXATIONAS/BI
E11 1 RELAXATIONAT/BI
E12 1 RELAXATIONATION/BI

=> s e3 or e8

216874 RELAXATION/BI
2096 RELAXATIONAL/BI
L12 217668 RELAXATION/BI OR RELAXATIONAL/BI

=> e narcotic

E1 1 NARCOTEINE/BI
E2 1 NARCOTHERAPY/BI
E3 7308 --> NARCOTIC/BI
E4 2 NARCOTICA/BI
E5 7 NARCOTICALLY/BI
E6 1 NARCOTICANTAGONIST/BI
E7 1 NARCOTICDOSE/BI
E8 1 NARCOTICI/BI
E9 1 NARCOTICISM/BI
E10 6 NARCOTICIZED/BI
E11 2 NARCOTICLIKE/BI
E12 5274 NARCOTICS/BI

=> a e3 or e12

A IS NOT A RECOGNIZED COMMAND

The previous command name entered was not recognized by the system.
For a list of commands available to you in the current file, enter
"HELP COMMANDS" at an arrow prompt (=>).

=> s e3 or e12

7308 NARCOTIC/BI
5274 NARCOTICS/BI
L13 10398 NARCOTIC/BI OR NARCOTICS/BI

=> e hypnotic

E1 1 HYPNOTHERAPEUTIC/BI
E2 7 HYPNOTHERAPY/BI
E3 5466 --> HYPNOTIC/BI
E4 1 HYPNOTICA/BI
E5 1 HYPNOTICACTIVITY/BI
E6 22 HYPNOTICALLY/BI
E7 1 HYPNOTICATES/BI
E8 1 HYPNOTICLIKE/BI
E9 1 HYPNOTICO/BI
E10 6371 HYPNOTICS/BI

E11 1 HYPNOTICSEDATIVES/BI
E12 6 HYPNOTIES/BI

=> s e3 or e10
5466 HYPNOTIC/BI
6371 HYPNOTICS/BI
L14 9427 HYPNOTIC/BI OR HYPNOTICS/BI

=> e somulence
E1 1 SOMUKI/BI
E2 1 SOMULA/BI
E3 0 --> SOMULENCE/BI
E4 2 SOMUNCURA/BI
E5 1 SOMUNCURENSIS/BI
E6 1 SOMUNCURICA/BI
E7 1 SOMUNIFERUM/BI
E8 1 SOMUOLENCE/BI
E9 2 SOMV/BI
E10 1 SOMVARSKY/BI
E11 1 SOMVIX/BI
E12 3 SOMW/BI

=> e somul
E1 2 SOMUCH/BI
E2 1 SOMUKI/BI
E3 0 --> SOMUL/BI
E4 1 SOMULA/BI
E5 2 SOMUNCURA/BI
E6 1 SOMUNCURENSIS/BI
E7 1 SOMUNCURICA/BI
E8 1 SOMUNIFERUM/BI
E9 1 SOMUOLENCE/BI
E10 2 SOMV/BI
E11 1 SOMVARSKY/BI
E12 1 SOMVIX/BI

=> e somnia
E1 1 SOMNHCOCF3/BI
E2 1 SOMNI/BI
E3 1 --> SOMNIA/BI
E4 2 SOMNIANS/BI
E5 3 SOMNIARIA/BI
E6 2 SOMNIF/BI
E7 15 SOMNIFACIENT/BI
E8 14 SOMNIFACIENTS/BI
E9 5 SOMNIFAINE/BI
E10 1 SOMNIFEA/BI
E11 84 SOMNIFEN/BI
E12 10 SOMNIFENE/BI

=> e insomnia
E1 1 INSOMMIA/BI
E2 1 INSOMMNIA/BI
E3 1393 --> INSOMNIA/BI
E4 54 INSOMNIAC/BI
E5 82 INSOMNIACS/BI
E6 5 INSOMNIAS/BI
E7 3 INSOMNIC/BI
E8 1 INSOMNIOUS/BI
E9 1 INSOMORPHOUS/BI
E10 8 INSOMUCH/BI
E11 2 INSON/BI

E12 2 INSONATE/BI

=> s e3-e8

1393 INSOMNIA/BI
54 INSOMNIAC/BI
82 INSOMNIACS/BI
5 INSOMNIAS/BI
3 INSOMNIC/BI
1 INSOMNIOUS/BI

L15 1431 (INSOMNIA/BI OR INSOMNIAC/BI OR INSOMNIACS/BI OR INSOMNIAS/BI
OR INSOMNIC/BI OR INSOMNIOUS/BI)

=> s l1

L16 717 L1

=> s l2

L17 214 L2

=> s l3

L18 350 L3

=> s l4

L19 1195 L4

=> s l5

L20 80 L5

=> s l6

L21 487 L6

=> s l8

L22 621 L8

=> s l11

9549 SEDATIVE/BI
1 SEDATIVEACTION/BI
1 SEDATIVEFOR/BI
4 SEDATIVEHYPNOTIC/BI
3 SEDATIVELIKE/BI
3 SEDATIVELY/BI
7525 SEDATIVES/BI

L23 12852 (SEDATIVE/BI OR SEDATIVEACTION/BI OR SEDATIVEFOR/BI OR SEDATIVEH
YPNOTIC/BI OR SEDATIVELIKE/BI OR SEDATIVELY/BI OR SEDATIVES/BI)

=> s l11

9549 SEDATIVE/BI
1 SEDATIVEACTION/BI
1 SEDATIVEFOR/BI
4 SEDATIVEHYPNOTIC/BI
3 SEDATIVELIKE/BI
3 SEDATIVELY/BI
7525 SEDATIVES/BI

L24 12852 (SEDATIVE/BI OR SEDATIVEACTION/BI OR SEDATIVEFOR/BI OR SEDATIVEH
YPNOTIC/BI OR SEDATIVELIKE/BI OR SEDATIVELY/BI OR SEDATIVES/BI)

=> s l9

L25 422 L9

=> s 16 and l10

487 L6

L26 1 L6 AND L10

=> d 126 1 all

L26 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian
CC 1-11 (Pharmacology)
AB In expts. on male mice and rats some behavioral effects of the diterpene sclareol glucol (SG) were studied. SG in higher doses (50, 100 mg/kg) enhanced the sensitivity to touch and pain, shortened the duration of hexobarbital **sleep**, stimulated exploratory behavior in rats in open field, exerted no anti-convulsive effect and increased the mortality from pentylenetetrazole seizures. In higher doses (300, 500, and 1000 mg/kg) SG induced seizures.
ST sclareol glucol behavior nervous system
IT Behavior
Nervous system
(sclareol glucol effect on)
IT 38419-75-9
RL: PRP (Properties)
(behavioral and nervous systems effects of)

=> FIL REGISTRY

| COST IN U.S. DOLLARS | SINCE FILE ENTRY | TOTAL SESSION |
|--|------------------|---------------|
| FULL ESTIMATED COST | 67.11 | 132.82 |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE ENTRY | TOTAL SESSION |
| CA SUBSCRIBER PRICE | -0.65 | -0.65 |

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 14 JUL 2003 HIGHEST RN 548428-18-8
DICTIONARY FILE UPDATES: 14 JUL 2003 HIGHEST RN 548428-18-8

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details:
<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

=> S 38419-75-9/RN

L27 1 38419-75-9/RN

=> SET NOTICE 1 DISPLAY

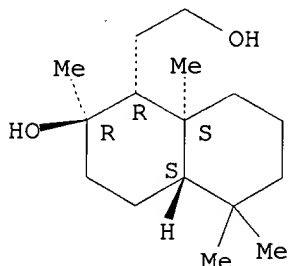
NOTICE SET TO 1 U.S. DOLLAR FOR DISPLAY COMMAND
SET COMMAND COMPLETED

=> D L27 SQIDE 1-

YOU HAVE REQUESTED DATA FROM 1 ANSWERS - CONTINUE? Y/(N):y
THE ESTIMATED COST FOR THIS REQUEST IS 5.63 U.S. DOLLARS
DO YOU WANT TO CONTINUE WITH THIS REQUEST? (Y)/N:y

L27 ANSWER 1 OF 1 REGISTRY COPYRIGHT 2003 ACS
RN 38419-75-9 REGISTRY
CN 1-Naphthaleneethanol, decahydro-2-hydroxy-2,5,5,8a-tetramethyl-,
(1R,2R,4aS,8aS)- (9CI) (CA INDEX NAME)
OTHER CA INDEX NAMES:
CN 1-Naphthaleneethanol, decahydro-2-hydroxy-2,5,5,8a-tetramethyl-,
[1R-(1.alpha.,2.beta.,4a.beta.,8a.alpha.)]-
OTHER NAMES:
CN 13,14,15,16-Tetranorlabdane-8,12-diol
CN 13,14,15,16-Tetranorlabdane-8.alpha.,12-diol
CN Ambroxdiol
CN AT 1
CN Sclareol glycol
FS STEREOSEARCH
MF C16 H30 O2
LC STN Files: BEILSTEIN*, BIOSIS, CA, CAPLUS, CASREACT, CHEMINFORMRX, DDFU,
DRUGU, TOXCENTER, USPATFULL
(*File contains numerically searchable property data)

Absolute stereochemistry. Rotation (-).



PROPERTY DATA AVAILABLE IN THE 'PROP' FORMAT

84 REFERENCES IN FILE CA (1957 TO DATE)
84 REFERENCES IN FILE CAPLUS (1957 TO DATE)

=> SET NOTICE LOGIN DISPLAY

NOTICE SET TO OFF FOR DISPLAY COMMAND
SET COMMAND COMPLETED

=>

| | | | |
|--|--|------------|---------|
| => file caplus | | SINCE FILE | TOTAL |
| COST IN U.S. DOLLARS | | ENTRY | SESSION |
| FULL ESTIMATED COST | | 2.08 | 134.90 |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | | SINCE FILE | TOTAL |
| CA SUBSCRIBER PRICE | | ENTRY | SESSION |
| | | 0.00 | -0.65 |

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FILE COVERS 1907 - 15 Jul 2003 VOL 139 ISS 3
 FILE LAST UPDATED: 14 Jul 2003 (20030714/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

| | |
|----|----------------|
| L1 | 43 S CEDROL |
| L2 | 33 S PATCHOULI |
| L3 | 36 S SANTALOL |
| L4 | 31 S BISABOLOL |
| | E BISABOLOL |
| L5 | 2 S VETIVEROL |
| L6 | 30 S SCLAREOL |
| L7 | 0 S GLOBUOL |
| L8 | 9 S GLOBULOL |
| L9 | 11 S GUAIOL |

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

E SLEEP

| | |
|-----|-------------------|
| L10 | 15616 S E3 |
| | E SEDATIVE |
| L11 | 12852 S E3-E9 |
| | E RELAXATION |
| L12 | 217668 S E3 OR E8 |
| | E NARCOTIC |
| L13 | 10398 S E3 OR E12 |
| | E HYPNOTIC |
| L14 | 9427 S E3 OR E10 |
| | E SOMULENCE |
| | E SOMUL |
| | E SOMNIA |

E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3
L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8
L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
L27 1 S 38419-75-9/RN
SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003

=> s 110 and 116
L28 2 L10 AND L16

=> d 128 1-2

L28 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS
AN 2003:132334 CAPLUS
DN 138:158861
TI Sleep-inducing dentifrices containing menthol and cedrene sesquiterpene alcohols
IN Itano, Morihide; Oshino, Kazushi; Nagashima, Yoshinao; Yata, Sachihiro
PA Kao Corp., Japan
SO Jpn. Kokai Tokkyo Koho, 6 pp.
CODEN: JKXXAF
DT Patent
LA Japanese
FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------------|------|----------|-----------------|----------|
| ----- | --- | ----- | ----- | ----- |
| PI JP 2003048827 | A2 | 20030221 | JP 2001-234832 | 20010802 |
| PRAI JP 2001-234832 | | 20010802 | | |

L28 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS
AN 2001:597788 CAPLUS
DN 135:170507
TI Autonomic-controlling agents containing sesquiterpene alcohols
IN Nagashima, Yoshinao; Sugata, Keiichi; Yada, Yukihiro; Fukuda, Kazuyuki
PA Kao Corp., Japan
SO PCT Int. Appl., 48 pp.
CODEN: PIXXD2
DT Patent
LA Japanese
FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|-----------------|----------|
| ----- | --- | ----- | ----- | ----- |
| PI WO 2001058435 | A1 | 20010816 | WO 2001-JP928 | 20010209 |
| W: JP, US | | | | |
| RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, | | | | |
| PT, SE, TR | | | | |
| EP 1170005 | A1 | 20020109 | EP 2001-902822 | 20010209 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, | | | | |

IE, FI
US 2002151600 A1 20021017 US 2001-972887 20011010
PRAI JP 2000-38260 A 20000210
WO 2001-JP928 W 20010209

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s l16 an l11

MISSING OPERATOR L16 AN

The search profile that was entered contains terms or
nested terms that are not separated by a logical operator.

=> s l16 and l11

L29 1 L16 AND L11

=> d 129

L29 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 2001:597788 CAPLUS

DN 135:170507

TI Autonomic-controlling agents containing sesquiterpene alcohols

IN Nagashima, Yoshinao; Sugata, Keiichi; Yada, Yukihiro; Fukuda, Kazuyuki

PA Kao Corp., Japan

SO PCT Int. Appl., 48 pp.

CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|---|------|----------|-----------------|----------|
| PI | WO 2001058435 | A1 | 20010816 | WO 2001-JP928 | 20010209 |
| | W: JP, US | | | | |
| | RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, | | | | |
| | PT, SE, TR | | | | |
| | EP 1170005 | A1 | 20020109 | EP 2001-902822 | 20010209 |
| | R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, | | | | |
| | IE, FI | | | | |
| | US 2002151600 | A1 | 20021017 | US 2001-972887 | 20011010 |
| PRAI | JP 2000-38260 | A | 20000210 | | |
| | WO 2001-JP928 | W | 20010209 | | |

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s l16 and l12

L30 0 L16 AND L12

=> s l16 and l13

L31 0 L16 AND L13

=> s l16 and l14

MISSING OPERATOR L16 AND L14

The search profile that was entered contains terms or
nested terms that are not separated by a logical operator.

=> s l16 and l14

L32 1 L16 AND L14

=> d 132

L32 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS

AN 2001:597788 CAPLUS
DN 135:170507
TI Autonomic-controlling agents containing sesquiterpene alcohols
IN Nagashima, Yoshinao; Sugata, Keiichi; Yada, Yukihiro; Fukuda, Kazuyuki
PA Kao Corp., Japan
SO PCT Int. Appl., 48 pp.
CODEN: PIXXD2

DT Patent

LA Japanese

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|---|------|----------|-----------------|----------|
| PI | WO 2001058435 | A1 | 20010816 | WO 2001-JP928 | 20010209 |
| | W: JP, US | | | | |
| | RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, | | | | |
| | PT, SE, TR | | | | |
| | EP 1170005 | A1 | 20020109 | EP 2001-902822 | 20010209 |
| | R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, | | | | |
| | IE, FI | | | | |
| | US 2002151600 | A1 | 20021017 | US 2001-972887 | 20011010 |
| PRAI | JP 2000-38260 | A | 20000210 | | |
| | WO 2001-JP928 | W | 20010209 | | |

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s l15 and l16
L33 0 L15 AND L16

=> s l17 and l10
L34 0 L17 AND L10

=> s l17 and l11
L35 0 L17 AND L11

=> s l17 and l13
L36 0 L17 AND L13

=> s l17 and l15
L37 0 L17 AND L15

=> s l18 and l10
L38 0 L18 AND L10

=> s l18 and l13
L39 0 L18 AND L13

=> s l19 and l10
L40 0 L19 AND L10

=> s l118 and l12
L41 0 LL18
0 LL18 AND L12

=> s l18 and l12
L42 0 L18 AND L12

=> s l19 and l12
L43 1 L19 AND L12

=> d 143

L43 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1993:616760 CAPLUS
DN 119:216760
TI Calcium antagonistic properties of the sesquiterpene T-cadinol and related substances: structure-activity studies
AU Zygmunt, P. M.; Larsson, B.; Sterner, O.; Vinge, E.; Hoegestaett, E. D.
CS Dep. Clin. Pharmacol., Univ. Hosp. Lund, Lund, S-22185, Swed.
SO Pharmacology & Toxicology (Oxford, United Kingdom) (1993), 73(1), 3-9
CODEN: PHTOEH; ISSN: 0901-9928
DT Journal
LA English

=> s 43 1 all
215843 43
7600426 1
1664927 ALL
L44 1 43 1 ALL
(43(W) 1(W)ALL)

=> d 143 1 all

L43 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1993:616760 CAPLUS
DN 119:216760
TI Calcium antagonistic properties of the sesquiterpene T-cadinol and related substances: structure-activity studies
AU Zygmunt, P. M.; Larsson, B.; Sterner, O.; Vinge, E.; Hoegestaett, E. D.
CS Dep. Clin. Pharmacol., Univ. Hosp. Lund, Lund, S-22185, Swed.
SO Pharmacology & Toxicology (Oxford, United Kingdom) (1993), 73(1), 3-9
CODEN: PHTOEH; ISSN: 0901-9928
DT Journal
LA English
CC 1-3 (Pharmacology)
AB The calcium antagonistic properties of (+)-T-cadinol, some of its stereoisomers and related terpenes were investigated in both functional and radioligand binding studies, and the effects were compared with those of the dihydropyridine calcium antagonist (.-+.-)-nimodipine. In the isolated rat aorta, the terpenes relaxed contractions induced by 60 mM K⁺ more potently than those induced by phenylephrine. (+)-T-cadinol and its stereoisomers were the most potent among the terpenes to relax K⁺-induced contractions, whereas they were approx. 10,000 times less potent than (.-+.-)-nimodipine in this regard. Binding of the dihydropyridine radioligand [3H]-(+)-PN200-110 was studied on rat cerebral cortical membranes. Displacement and satn. studies indicated that (+)-T-cadinol caused a competitive inhibition of binding. The log Ki values for (+)-T-cadinol and (.-+.-)-nimodipine from displacement studies (-4.7 and -9.2) corresponded with the log RC50 values for relaxation of K⁺-contracted rat aortas (-5.0 and -9.0). For the terpenes, there was a significant correlation ($P < 0.001$, $r_s = 0.89$) between displacement of dihydropyridine binding and the ability to relax K⁺-induced contractions. The structures of three terpenes were chem. modified by blocking hydroxyl groups. The potency of these derivs., as well as the naturally occurring deriv. 2-oxo-T-cadinol, to relax K⁺-induced contractions was not correlated to the lipophilicity of the compds. Instead, other qualities appear to be of importance for the functional effects. The authors' results suggest that (+)-T-cadinol and related terpenes may represent a new chem. class of calcium antagonists, which interact with dihydropyridine binding sites on the voltage-operated calcium channels.
ST calcium antagonist terpene T cadinol structure
IT Terpenes and Terpenoids, biological studies
RL: BIOL (Biological study)

(calcium antagonism by, structure in relation to)
IT Lipophilicity
(of sesquiterpene T-cadinol and related substances, calcium antagonism
in relation to)
IT Ion channel blockers
(calcium, sesquiterpene T-cadinol and related substances as, structure
in relation to)
IT Molecular structure-biological activity relationship
(calcium channel-blocking, of sesquiterpene T-cadinol and related
substances)
IT Receptors
RL: BIOL (Biological study)
(dihydropyridine, sesquiterpene T-cadinol and related substances
binding to, calcium antagonism by, structure in relation to)
IT 481-34-5, (-)-alpha.-Cadinol 2216-51-5, (-)-Menthol 5937-11-1,
(+)-T-Cadinol 19435-97-3 19912-62-0, (-)-T-Muurolol **23089-26-1**
, (-)-alpha.-Bisabolol 53402-16-7 74638-12-3, (-)-Furosardonin A
129058-89-5, (-)-Tremediol 150718-45-9 150718-46-0 150718-47-1
RL: BIOL (Biological study)
(calcium antagonism by, structure in relation to)

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

L1 43 S CEDROL
L2 33 S PATCHOULI
L3 36 S SANTALOL
L4 31 S BISABOLOL
E BISABOLOL
L5 2 S VETIVEROL
L6 30 S SCLAREOL
L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

E SLEEP
L10 15616 S E3
E SEDATIVE
L11 12852 S E3-E9
E RELAXATION
L12 217668 S E3 OR E8
E NARCOTIC
L13 10398 S E3 OR E12
E HYPNOTIC
L14 9427 S E3 OR E10
E SOMULENCE
E SOMUL
E SOMNIA
E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3
L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8
L23 12852 S L11

L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
L27 1 S 38419-75-9/RN
SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003
L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16
L34 0 S L17 AND L10
L35 0 S L17 AND L11
L36 0 S L17 AND L13
L37 0 S L17 AND L15
L38 0 S L18 AND L10
L39 0 S L18 AND L13
L40 0 S L19 AND L10
L41 0 S LL18 AND L12
L42 0 S L18 AND L12
L43 1 S L19 AND L12
L44 1 S 43 1 ALL

=> s 119 and 115
L45 0 L19 AND L15

=> s 120 and 110
L46 0 L20 AND L10

=> s 120 and 112
L47 0 L20 AND L12

=> s 121 and 110
L48 1 L21 AND L10

=> d 148

L48 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian

=> d 148 all

L48 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.

SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian
CC 1-11 (Pharmacology)
AB In expts. on male mice and rats some behavioral effects of the diterpene sclareol glucol (SG) were studied. SG in higher doses (50, 100 mg/kg) enhanced the sensitivity to touch and pain, shortened the duration of hexobarbital **sleep**, stimulated exploratory behavior in rats in open field, exerted no anti-convulsive effect and increased the mortality from pentylenetetrazole seizures. In higher doses (300, 500, and 1000 mg/kg) SG induced seizures.
ST sclareol glucol behavior nervous system
IT Behavior
Nervous system
(sclareol glucol effect on)
IT 38419-75-9
RL: PRP (Properties)
(behavioral and nervous systems effects of)

=> s 122 and 112
L49 0 L22 AND L12

=> s 125 and 110
L50 0 L25 AND L10

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

L1 43 S CEDROL
L2 33 S PATCHOULI
L3 36 S SANTALOL
L4 31 S BISABOLOL
E BISABOLOL
L5 2 S VETIVEROL
L6 30 S SCLAREOL
L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

E SLEEP
L10 15616 S E3
E SEDATIVE
L11 12852 S E3-E9
E RELAXATION
L12 217668 S E3 OR E8
E NARCOTIC
L13 10398 S E3 OR E12
E HYPNOTIC
L14 9427 S E3 OR E10
E SOMULENCE
E SOMUL
E SOMNIA
E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3

L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8
L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
L27 1 S 38419-75-9/RN
SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003
L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16
L34 0 S L17 AND L10
L35 0 S L17 AND L11
L36 0 S L17 AND L13
L37 0 S L17 AND L15
L38 0 S L18 AND L10
L39 0 S L18 AND L13
L40 0 S L19 AND L10
L41 0 S LL18 AND L12
L42 0 S L18 AND L12
L43 1 S L19 AND L12
L44 1 S 43 1 ALL
L45 0 S L19 AND L15
L46 0 S L20 AND L10
L47 0 S L20 AND L12
L48 1 S L21 AND L10
L49 0 S L22 AND L12
L50 0 S L25 AND L10

=> e nervious
E1 1 NERVIOSA/BI
E2 1 NERVIOSO/BI
E3 3 --> NERVIOUS/BI
E4 1 NERVIS/BI
E5 6 NERVISTEROL/BI
E6 1 NERVIUM/BI
E7 1 NERVNATA/BI
E8 5 NERVNAYA/BI
E9 1 NERVNNOE/BI
E10 2 NERVNOGO/BI
E11 29 NERVNNOI/BI
E12 1 NERVNOMYSHECHNOGO/BI

=> e nervous
E1 2 NERVOUR/BI
E2 1 NERVOURSE/BI
E3 163958 --> NERVOUS/BI
E4 1 NERVOUSDEPRESSANT/BI
E5 4 NERVOUSE/BI
E6 27 NERVOUSLY/BI
E7 254 NERVOUSNESS/BI
E8 1 NERVOUSSVSTEM/BI

E9 7 NERVOUSSYSTEM/BI
E10 1 NERVOOUS/BI
E11 1 NERVOUW/BI
E12 2 NERVOV/BI

=> s e3-e7

163958 NERVOUS/BI
1 NERVOUSDEPRESSANT/BI
4 NERVOUSE/BI
27 NERVOUSLY/BI
254 NERVOUSNESS/BI

L51 164186 (NERVOUS/BI OR NERVOUSDEPRESSANT/BI OR NERVOUSE/BI OR NERVOUSLY/BI OR NERVOUSNESS/BI)

=> s l51 and l16

L52 1 L51 AND L16

=> d 152 all

L52 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 2001:597788 CAPLUS
DN 135:170507
TI Autonomic-controlling agents containing sesquiterpene alcohols
IN Nagashima, Yoshinao; Sugata, Keiichi; Yada, Yukihiro; Fukuda, Kazuyuki
PA Kao Corp., Japan
SO PCT Int. Appl., 48 pp.
CODEN: PIXXD2
DT Patent
LA Japanese
IC ICM A61K031-045
ICS A61K007-46; A61P025-02; A61P025-20
CC 62-5 (Essential Oils and Cosmetics)
Section cross-reference(s): 17, 63
FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---|------|----------|-----------------|----------|
| WO 2001058435 | A1 | 20010816 | WO 2001-JP928 | 20010209 |
| W: JP, US | | | | |
| RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, | | | | |
| PT, SE, TR | | | | |
| EP 1170005 | A1 | 20020109 | EP 2001-902822 | 20010209 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, | | | | |
| IE, FI | | | | |
| US 2002151600 | A1 | 20021017 | US 2001-972887 | 20011010 |
| PRAI JP 2000-38260 | A | 20000210 | | |
| WO 2001-JP928 | W | 20010209 | | |

AB Disclosed are autonomic-controlling agents exerting sedative, sleep-inducing, and stress-relieving effects on humans regardless of differences among individuals in the sensitivity or preference to smell. These agents contain as the main active ingredient sesquiterpene alcs. having a b.p. of .gtoreq. 250.degree. under atm. pressure, in particular, cedrol.

ST sesquiterpene alc autonomic control sedative; cedrol hypnotic stress relief aroma therapy

IT Hypnotics and Sedatives
(autonomic-controlling agents contg. sesquiterpene alcs.)

IT Candy
(autonomic-controlling agents contg. sesquiterpene alcs. in)

IT Nervous system
(autonomic; autonomic-controlling agents contg. sesquiterpene alcs.)

IT Cosmetics
(creams, massage; autonomic-controlling agents contg. sesquiterpene

alcs. in)
IT Medical goods
(face masks contg. cedrol; autonomic-controlling agents contg.
sesquiterpene alcs. in)
IT Sesquiterpenes
RL: BUU (Biological use, unclassified); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(hydroxy; autonomic-controlling agents contg. sesquiterpene alcs.)
IT Stress, animal
(relief; autonomic-controlling agents contg. sesquiterpene alcs.)
IT Odor and Odorous substances
(therapy; autonomic-controlling agents contg. sesquiterpene alcs.)
IT 77-53-2, Cedrol
RL: BUU (Biological use, unclassified); THU (Therapeutic use); BIOL
(Biological study); USES (Uses)
(autonomic-controlling agents contg. sesquiterpene alcs.)

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD

RE

- (1) American Chemical Society; Database CAPLUS on STN
- (2) American Chemical Society; Database CAPLUS on STN
- (3) American Chemical Society; Database CAPLUS on STN
- (4) American Chemical Society; Database CAPLUS on STN
- (5) International Flavors And Fragrances Inc; US 4670264 A CAPLUS
- (6) International Flavors And Fragrances Inc; US 4670463 A CAPLUS
- (7) International Flavors And Fragrances Inc; US 4671959 A CAPLUS
- (8) International Flavors And Fragrances Inc; JP 61267526 A CAPLUS
- (9) International Flavors And Fragrances Inc; EP 183436 A2 1986 CAPLUS
- (10) Kobayashi Pharmaceutical Co Ltd; JP 1025245 A 1998
- (11) Narisu Keshohin K K; JP 11343497 A 1999 CAPLUS
- (12) Sawada, K; Nippon Aji to Nioi Gakkaishi 1999, V6(3), P465 CAPLUS

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

L1 43 S CEDROL
L2 33 S PATCHOULI
L3 36 S SANTALOL
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L5 2 S VETIVEROL
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L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

E SLEEP
L10 15616 S E3
E SEDATIVE
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E RELAXATION
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E NARCOTIC
L13 10398 S E3 OR E12
E HYPNOTIC
L14 9427 S E3 OR E10
E SOMULENCE
E SOMUL
E SOMNIA

E INSOMNIA
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 L18 350 S L3
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 L20 80 S L5
 L21 487 S L6
 L22 621 S L8
 L23 12852 S L11
 L24 12852 S L11
 L25 422 S L9
 L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
 L27 1 S 38419-75-9/RN
 SET NOTICE 1 DISPLAY
 SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003
 L28 2 S L10 AND L16
 L29 1 S L16 AND L11
 L30 0 S L16 AND L12
 L31 0 S L16 AND L13
 L32 1 S L16 AND L14
 L33 0 S L15 AND L16
 L34 0 S L17 AND L10
 L35 0 S L17 AND L11
 L36 0 S L17 AND L13
 L37 0 S L17 AND L15
 L38 0 S L18 AND L10
 L39 0 S L18 AND L13
 L40 0 S L19 AND L10
 L41 0 S L20 AND L12
 L42 0 S L18 AND L12
 L43 1 S L19 AND L12
 L44 1 S 43 1 ALL
 L45 0 S L19 AND L15
 L46 0 S L20 AND L10
 L47 0 S L20 AND L12
 L48 1 S L21 AND L10
 L49 0 S L22 AND L12
 L50 0 S L25 AND L10
 E NERVOUS
 E NERVOUS
 L51 164186 S E3-E7
 L52 1 S L51 AND L16

| | | | |
|--|------------------|---------------|--|
| => file reg | | | |
| COST IN U.S. DOLLARS | SINCE FILE ENTRY | TOTAL SESSION | |
| FULL ESTIMATED COST | 38.04 | 172.94 | |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE ENTRY | TOTAL SESSION | |
| CA SUBSCRIBER PRICE | -1.95 | -2.60 | |

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Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 14 JUL 2003 HIGHEST RN 548428-18-8
DICTIONARY FILE UPDATES: 14 JUL 2003 HIGHEST RN 548428-18-8

TSCA INFORMATION NOW CURRENT THROUGH JANUARY 6, 2003

Please note that search-term pricing does apply when conducting SmartSELECT searches.

Crossover limits have been increased. See HELP CROSSOVER for details.

Experimental and calculated property data are now available. See HELP PROPERTIES for more information. See STNote 27, Searching Properties in the CAS Registry File, for complete details:

<http://www.cas.org/ONLINE/STN/STNOTES/stnotes27.pdf>

=> e farnesol

| | | |
|-----|--------|------------------------|
| E1 | 6 | FARNESOIC/BI |
| E2 | 29 | FARNESOID/BI |
| E3 | 75 --> | FARNESOL/BI |
| E4 | 1 | FARNESOLATE/BI |
| E5 | 1 | FARNESOLIC/BI |
| E6 | 1 | FARNESONE/BI |
| E7 | 1 | FARNESONITRILE/BI |
| E8 | 1 | FARNESOXY/BI |
| E9 | 5 | FARNESOYL/BI |
| E10 | 1 | FARNESOYLHYDRO/BI |
| E11 | 1 | FARNESOYLHYDROXAMIC/BI |
| E12 | 2 | FARNESOYLPENICILL/BI |

=> s e3

| | | |
|-----|----|-------------|
| L53 | 75 | FARNESOL/BI |
|-----|----|-------------|

=> e eugenol

| | | |
|-----|---------|--------------------|
| E1 | 2 | EUGENODIL/BI |
| E2 | 2 | EUGENODILOL/BI |
| E3 | 165 --> | EUGENOL/BI |
| E4 | 1 | EUGENOLATE/BI |
| E5 | 1 | EUGENOLATO/BI |
| E6 | 1 | EUGENOLGLYC/BI |
| E7 | 1 | EUGENOLGLYCOL/BI |
| E8 | 1 | EUGENOLGLYCOLIC/BI |
| E9 | 1 | EUGENOLOL/BI |
| E10 | 1 | EUGENON/BI |
| E11 | 1 | EUGENONE/BI |
| E12 | 1 | EUGENOXIDE/BI |

=> s e3

| | | |
|-----|-----|------------|
| L54 | 165 | EUGENOL/BI |
|-----|-----|------------|

=> s geranyl linalool

| | | |
|------|----------|----------------------|
| 1021 | GERANYL | |
| 92 | LINALOOL | |
| L55 | 4 | GERANYL LINALOOL |
| | | (GERANYL(W)LINALOOL) |

=> e cedrenol

| | | |
|----|-------|------------------------|
| E1 | 1 | CEDRENEDICARBOXYLIC/BI |
| E2 | 1 | CEDRENIC/BI |
| E3 | 9 --> | CEDRENOL/BI |

E4 2 CEDRENON/BI
E5 2 CEDRENONE/BI
E6 1 CEDRENYL/BI
E7 1 CEDRI/BI
E8 1 CEDRIC/BI
E9 8 CEDRIN/BI
E10 1 CEDRINOSIDE/BI
E11 1 CEDRIRET/BI
E12 7 CEDRO/BI

=> s e3
L56 9 CEDRENOL/BI

=> e isopytol
E1 1 ISOPYTHALDINE/BI
E2 1 ISOPYTHALINE/BI
E3 0 --> ISOPYTOL/BI
E4 1 ISOQIN/BI
E5 1 ISOQINOL/BI
E6 1 ISOQINOLINE/BI
E7 1 ISOQU/BI
E8 2 ISOQUADR/BI
E9 2 ISOQUADRONE/BI
E10 2 ISOQUASSIN/BI
E11 2 ISOQUASSINIC/BI
E12 1 ISOQUATER/BI

=> e isophytol
E1 10 ISOPHYT/BI
E2 1 ISOPHYTO/BI
E3 6 --> ISOPHYTOL/BI
E4 1 ISOPHYTOLACCAGENIN/BI
E5 1 ISOPHYTOLACCINIC/BI
E6 1 ISOPHYTOSPHINGO/BI
E7 1 ISOPHYTOSPHINGOSINE/BI
E8 4 ISOPHYTYL/BI
E9 4 ISOPI/BI
E10 2 ISOPICHIEREN/BI
E11 2 ISOPICHIERENOL/BI
E12 1 ISOPICHIERENYL/BI

=> s e3
L57 6 ISOPHYTOL/BI

=> e nerolidol
E1 1 NEROLIDI/BI
E2 1 NEROLIDIOL/BI
E3 41 --> NEROLIDOL/BI
E4 8 NEROLIDYL/BI
E5 1 NEROLIDYLCAECH/BI
E6 1 NEROLIDYLCAECHOL/BI
E7 4 NEROLIN/BI
E8 1 NEROLINE/BI
E9 1 NEROLIT/BI
E10 2 NEROLOA/BI
E11 1 NERONE/BI
E12 1 NERONIN/BI

=> s e3
L58 41 NEROLIDOL/BI

=> file caplus

| COST IN U.S. DOLLARS | SINCE FILE ENTRY | TOTAL SESSION |
|--|------------------|---------------|
| FULL ESTIMATED COST | 32.74 | 205.68 |
| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE ENTRY | TOTAL SESSION |
| CA SUBSCRIBER PRICE | 0.00 | -2.60 |

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FILE COVERS 1907 - 15 Jul 2003 VOL 139 ISS 3
 FILE LAST UPDATED: 14 Jul 2003 (20030714/ED)

This file contains CAS Registry Numbers for easy and accurate substance identification.

```
=> s 156 and l10
      90 L56
L59      1 L56 AND L10
```

```
=> d 159 all
```

L59 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
 AN 2003:132334 CAPLUS
 DN 138:158861
 TI **Sleep**-inducing dentifrices containing menthol and cedrene sesquiterpene alcohols
 IN Itano, Morihide; Oshino, Kazushi; Nagashima, Yoshinao; Yata, Sachihiro
 PA Kao Corp., Japan
 SO Jpn. Kokai Tokkyo Koho, 6 pp.
 CODEN: JKXXAF
 DT Patent
 LA Japanese
 IC ICM A61K031-045
 ICS A61P025-20; A61P025-26
 CC 63-6 (Pharmaceuticals)
 Section cross-reference(s): 62
 FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------------|------|----------|-----------------|----------|
| PI JP 2003048827 | A2 | 20030221 | JP 2001-234832 | 20010802 |
| PRAI JP 2001-234832 | | 20010802 | | |

AB The dentifrices contain (A) menthol (I) and (B) cedrene sesquiterpene alcs. such as cedrol or cedrenol at (A)/(B) wt. ratio 1:0.01-10. The dentifrices show **sleep**-inducing effect because cedrene sesquiterpene alcs. suppress awakening effect of menthol. The cedrene sesquiterpene alcs. do not inhibit awakening effect of menthol in a

parasympathetic state such as a time just after awakening. A dentifrice contg. l-I 0.3, peppermint oil 0.2, spearmint oil 0.2, cedrol 0.004, sorbitol 30.0, glycerin 18.0, CaCO₃ 15.0, SiO₂ 7.5, Na lauryl sulfate 1.2, CM-cellulose 1.2, propylene glycol 0.5%, and H₂O balance significantly shortened time for falling asleep.

ST **sleep** inducing dentifrice menthol cedrene sesquiterpene alc; cedrol suppression menthol awakening effect **sleep** inducing dentifrice

IT Sesquiterpenes
RL: BSU (Biological study, unclassified); COS (Cosmetic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(hydroxy, cedrene; **sleep**-inducing dentifrices contg. menthol and cedrene sesquiterpene alcs. to suppress awakening effect of menthol)

IT Essential oils
RL: COS (Cosmetic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(peppermint, menthol-contg.; **sleep**-inducing dentifrices contg. menthol and cedrene sesquiterpene alcs. to suppress awakening effect of menthol)

IT Dentifrices
Human
Sleep
(**sleep**-inducing dentifrices contg. menthol and cedrene sesquiterpene alcs. to suppress awakening effect of menthol)

IT Essential oils
RL: COS (Cosmetic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(spearmint, menthol-contg.; **sleep**-inducing dentifrices contg. menthol and cedrene sesquiterpene alcs. to suppress awakening effect of menthol)

IT 77-53-2, Cedrol 28231-03-0, Cedrenol
RL: BSU (Biological study, unclassified); COS (Cosmetic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(**sleep**-inducing dentifrices contg. menthol and cedrene sesquiterpene alcs. to suppress awakening effect of menthol)

IT 1490-04-6, Menthol 2216-51-5
RL: COS (Cosmetic use); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
(**sleep**-inducing dentifrices contg. menthol and cedrene sesquiterpene alcs. to suppress awakening effect of menthol)

=> s 156 and 112
90 L56
L60 0 L56 AND L12

=> s 156 and 115
90 L56
L61 0 L56 AND L15

=> s 156
L62 90 L56

=> s ls 162 and 110
MISSING OPERATOR LS L62
The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

=> s 162 and 110
L63 1 L62 AND L10

=> d 163

L63 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 2003:132334 CAPLUS
DN 138:158861
TI **sleep-inducing dentifrices containing menthol and cedrene sesquiterpene alcohols**
IN Itano, Morihide; Oshino, Kazushi; Nagashima, Yoshinao; Yata, Sachihiro
PA Kao Corp., Japan
SO Jpn. Kokai Tokkyo Koho, 6 pp.
CODEN: JKXXAF
DT Patent
LA Japanese
FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------------|------|----------|-----------------|----------|
| JP 2003048827 | A2 | 20030221 | JP 2001-234832 | 20010802 |
| PRAI JP 2001-234832 | | 20010802 | | |

=> s 162 and 112

L64 0 L62 AND L12

=> s 153

L65 3275 L53

=> s 165 and 110

L66 1 L65 AND L10

=> d 166

L66 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1973:24110 CAPLUS
DN 78:24110
TI Farnesol, a psychosedative and spasmolytic compound
AU Binet, L.; Binet, P.; Miocque, M.; Morin, H.; Pechery, C.; Roux, M.;
Bernier, A.; Rinjard, P.; Godon, M.
CS Fac. Sci. Pharm. Biol., Paris, Fr.
SO Therapie (1972), 27(5), 893-905
CODEN: THERAP; ISSN: 0040-5957
DT Journal
LA French

=> d 166 1 all

L66 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1973:24110 CAPLUS
DN 78:24110
TI Farnesol, a psychosedative and spasmolytic compound
AU Binet, L.; Binet, P.; Miocque, M.; Morin, H.; Pechery, C.; Roux, M.;
Bernier, A.; Rinjard, P.; Godon, M.
CS Fac. Sci. Pharm. Biol., Paris, Fr.
SO Therapie (1972), 27(5), 893-905
CODEN: THERAP; ISSN: 0040-5957
DT Journal
LA French
CC 1-5 (Pharmacodynamics)
AB When given i.v. or orally at .geq.100 mg/kg, synthetic farnesol [4602-84-0] (contg. a mixt. of stereoisomers) was a psychosedative in mice and rats. Except at high doses, farnesol did not inhibit psychomotor reactions (defense reflex), but it lowered the response to

psychic stimuli such as curiosity and caffeine-induced excitation. Farnesol did not cause catalepsy, nor did it antagonize pentetetrazole-induced convulsion. It prolonged barbiturate **sleep** without itself being a hypnotic. Farnesol also had a musculotropic-type spasmolytic action on the isolated rat intestine and guinea pig sphincter of Oddi contracted by acetylcholine, BaCl₂, histamine, or serotonin.

ST farnesol sedative spasmolytic; muscle relaxant farnesol; tranquilizer farnesol

IT Muscle relaxants
Tranquilizers
(farnesol)

IT 58-08-2, biological studies
RL: BIOL (Biological study)
(excitation from, farnesol inhibition of)

IT **4602-84-0**
RL: BIOL (Biological study)
(sedative and spasmolytic)

IT 76-74-4
RL: BIOL (Biological study)
(**sleep** from, farnesol potentiation of)

=> s 165 and 112
L67 9 L65 AND L12

=> d 167 1-9

L67 ANSWER 1 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 2001:459531 CAPLUS
DN 135:368270
TI Membrane properties of sodium 2- and 6-(poly)prenyl-substituted polyprenyl phosphates
AU Takajo, Saho; Nagano, Hajime; Dannenmuller, Olivier; Ghosh, Sangita; Marie Albrecht, Anne; Nakatani, Yoichi; Ourisson, Guy
CS Department of Chemistry, Faculty of Science, Ochanomizu University, Otsuka, Bunkyo-ku, Tokyo, 112-8610, Japan
SO New Journal of Chemistry (2001), 25(7), 917-929
CODEN: NJCHE5; ISSN: 1144-0546
PB Royal Society of Chemistry
DT Journal
LA English
RE.CNT 72 THERE ARE 72 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L67 ANSWER 2 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1998:511778 CAPLUS
DN 129:254789
TI Improvement of nitrergic **relaxation** by farnesol of the sphincter of Oddi from hypercholesterolemic rabbits
AU Szilvassy, Zoltan; Sari, Reka; Nemeth, Jozsef; Nagy, Istvan; Csati, Sandor; Lonovics, Janos
CS 1st Department Medicine, Albert Szent-Gyorgyi Medical University Szeged, Szeged, Hung.
SO European Journal of Pharmacology (1998), 353(1), 75-78
CODEN: EJPRAZ; ISSN: 0014-2999
PB Elsevier Science B.V.
DT Journal
LA English
RE.CNT 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L67 ANSWER 3 OF 9 CAPLUS COPYRIGHT 2003 ACS

AN 1995:711214 CAPLUS
DN 123:107919
TI Initiation of biosynthesis in cis polyisoprenes
AU Tanaka, Yasuyuki; Kawahara, Seiichi; Aik-Hwee, Eng; Shiba, Kenichi; Ohya, Norimasa
CS Fac. Technol., Tokyo Univ. Agric. Technol., Koganei, 184, Japan
SO Phytochemistry (1995), 39(4), 779-84
CODEN: PYTCAS; ISSN: 0031-9422
PB Elsevier
DT Journal
LA English

L67 ANSWER 4 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1995:704229 CAPLUS
DN 123:228534
TI Carbon-13 NMR study of farnesol, farnesyl acetate and farnesal stereoisomers: chemical shift assignment using lanthanide induced shifts
AU Bradesi, Pascale; Tomi, Felix; Casanova, Joseph
CS Lab. Helioenergetique, Univ. Corse, Ajaccio, 20000, Fr.
SO Canadian Journal of Applied Spectroscopy (1995), 40(3), 76-81
CODEN: CJSPEM; ISSN: 1183-7306
PB Polyscience Publications, Inc.
DT Journal
LA English

L67 ANSWER 5 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1995:15887 CAPLUS
DN 122:49876
TI Mechanism of the biosynthesis of farnesyl diphosphate. Changes in the structure of geranyl diphosphate during the chain elongation process
AU Hiraga, Y.; Ito, D. I.; Takano, T.; Sayo, T.; Ohta, S.; Suga, T.
CS Fac. Sci., Hiroshima Univ., Japan
SO Tennen Yuki Kagobutsu Toronkai Koen Yoshishu (1993), 35th, 337-44
CODEN: TYKYDS
DT Journal
LA English/Japanese

L67 ANSWER 6 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1990:174389 CAPLUS
DN 112:174389
TI Nuclear magnetic resonance studies of polyisoprenols in model membranes
AU Knudsen, Mark J.; Troy, Frederic A.
CS Sch. Med., Univ. California, Davis, CA, 95616, USA
SO Chemistry and Physics of Lipids (1989), 51(3-4), 205-12
CODEN: CPLIA4; ISSN: 0009-3084
DT Journal
LA English

L67 ANSWER 7 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1985:591806 CAPLUS
DN 103:191806
TI Deuterium NMR investigation of the organization and dynamics of polyisoprenols in membranes
AU De Ropp, Jeffrey S.; Troy, Frederic A.
CS Sch. Med., Univ. California, Davis, CA, 95616, USA
SO Journal of Biological Chemistry (1985), 260(29), 15669-74
CODEN: JBCHA3; ISSN: 0021-9258
DT Journal
LA English

L67 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1985:200582 CAPLUS

DN 102:200582
TI Direct detection of solanesol in tobacco by proton and carbon-13 magic angle spinning NMR
AU Wooten, Jan B.
CS Philip Morris Res. Cent., Richmond, VA, 23261, USA
SO Journal of Agricultural and Food Chemistry (1985), 33(3), 419-25
CODEN: JAFCAU; ISSN: 0021-8561
DT Journal
LA English

L67 ANSWER 9 OF 9 CAPLUS COPYRIGHT 2003 ACS
AN 1984:205310 CAPLUS
DN 100:205310
TI Chemical synthesis and deuterium NMR investigations of polyisoprenols: dynamics in model membranes
AU De Ropp, Jeffrey S.; Troy, Frederic A.
CS Sch. Med., Univ. California, Davis, CA, 95616, USA
SO Biochemistry (1984), 23(12), 2691-5
CODEN: BICHAW; ISSN: 0006-2960
DT Journal
LA English

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

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L2 33 S PATCHOULI
L3 36 S SANTALOL
L4 31 S BISABOLOL
E BISABOLOL
L5 2 S VETIVEROL
L6 30 S SCLAREOL
L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

E SLEEP
L10 15616 S E3
E SEDATIVE
L11 12852 S E3-E9
E RELAXATION
L12 217668 S E3 OR E8
E NARCOTIC
L13 10398 S E3 OR E12
E HYPNOTIC
L14 9427 S E3 OR E10
E SOMULENCE
E SOMUL
E SOMNIA
E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3
L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8

L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

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L27 1 S 38419-75-9/RN
SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003
L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16
L34 0 S L17 AND L10
L35 0 S L17 AND L11
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L39 0 S L18 AND L13
L40 0 S L19 AND L10
L41 0 S LL18 AND L12
L42 0 S L18 AND L12
L43 1 S L19 AND L12
L44 1 S 43 1 ALL
L45 0 S L19 AND L15
L46 0 S L20 AND L10
L47 0 S L20 AND L12
L48 1 S L21 AND L10
L49 0 S L22 AND L12
L50 0 S L25 AND L10
E NERVOUS
E NERVOUS
L51 164186 S E3-E7
L52 1 S L51 AND L16

FILE 'REGISTRY' ENTERED AT 16:16:22 ON 15 JUL 2003
E FARNESOL
L53 75 S E3
E EUGENOL
L54 165 S E3
L55 4 S GERANYL LINALOOL
E CEDRENOL
L56 9 S E3
E ISOPYTOL
E ISOPHYTOL
L57 6 S E3
E NEROLIDOL
L58 41 S E3

FILE 'CAPLUS' ENTERED AT 16:21:24 ON 15 JUL 2003
L59 1 S L56 AND L10
L60 0 S L56 AND L12
L61 0 S L56 AND L15
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L64 0 S L62 AND L12
L65 3275 S L53
L66 1 S L65 AND L10

L67 9 S L65 AND L12

=> s 165 and 113

L68 0 L65 AND L13

=> s 165 and 115

L69 0 L65 AND L15

=> s 154

L70 8961 L54

=> s 170 and 110

L71 8 L70 AND L10

=> d 171 1-8

L71 ANSWER 1 OF 8 CAPLUS COPYRIGHT 2003 ACS

AN 2002:555449 CAPLUS

DN 137:109483

TI Preparation of alanine 2,6-dialkoxyphenyl ester derivatives as hypnotics

IN Hamilton, Niall Morton; Bennett, David Jonathan

PA Akzo Nobel N.V., Neth.

SO PCT Int. Appl., 39 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----|---------------|--|----------|-----------------|--|
| PI | WO 2002057218 | A1 | 20020725 | WO 2002-EP994 | 20020117 |
| | W: | AE, AG, AL, AU, BA, BB, BG, BR, BZ, CA, CN, CO, CR, CU, CZ, DM, DZ, EC, EE, GD, GE, HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG, MK, MN, MX, MZ, NO, NZ, PH, PL, RO, RU, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM | | RW: | GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG |

PRAI EP 2001-200195 A 20010119

OS MARPAT 137:109483

RE.CNT 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L71 ANSWER 2 OF 8 CAPLUS COPYRIGHT 2003 ACS

AN 1998:169475 CAPLUS

DN 128:248580

TI Association of NO synthase inhibitors with trappers of reactive oxygen species

IN Chabrier De Lassauniere, Pierre-Etienne; Bigg, Denis

PA Societe De Conseils De Recherches Et D'applications Scientifiques (S.C.R.A.S, Fr.; Chabrier De Lassauniere, Pierre-Etienne; Bigg, Denis

SO PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DT Patent

LA French

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----|------------|---|----------|-----------------|----------|
| PI | WO 9809653 | A1 | 19980312 | WO 1997-FR1567 | 19970905 |
| | W: | AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, | | | |

PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG,
 US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
 RW: GH, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, DE, DK, ES, FI, FR,
 GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA,
 GN, ML, MR, NE, SN, TD, TG
 FR 2753098 A1 19980313 FR 1996-10875 19960906
 FR 2753098 B1 19981127
 AU 9742111 A1 19980326 AU 1997-42111 19970905
 AU 734296 B2 20010607
 EP 939654 A1 19990908 EP 1997-940183 19970905
 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
 IE, FI
 NZ 334597 A 20001027 NZ 1997-334597 19970905
 JP 2000517336 T2 20001226 JP 1998-512314 19970905
 RU 2174844 C2 20011020 RU 1999-106792 19970905
 US 6297281 B1 20011002 US 1999-254254 19990302
 NO 9901100 A 19990505 NO 1999-1100 19990305
 PRAI FR 1996-10875 A 19960906
 WO 1997-FR1567 W 19970905
 RE.CNT 1 THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L71 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2003 ACS
 AN 1996:287269 CAPLUS
 DN 125:1102
 TI Synthesis and pharmacological activity of a eugenol derivative
 AU Costa, J. A.; Oliveira, R. A. G.; Barbosa, J. M.; Souza Brito, A. R. M.
 CS Lab. Tecnologia Farmaceutica, UFPB, Joao Pessoa, Brazil
 SO Revista Brasileira de Farmacia (1994), 75(2), 40-5
 CODEN: RBFAAH; ISSN: 0370-372X
 PB Associacao Brasileira de Farmaceuticos
 DT Journal
 LA Portuguese

L71 ANSWER 4 OF 8 CAPLUS COPYRIGHT 2003 ACS
 AN 1990:491283 CAPLUS
 DN 113:91283
 TI Inhibition and induction of hepatic mixed function oxidase by
 phenylpropanoids from the seeds of *Myristica fragrans*
 AU Shin, Kuk Hyun; Woo, Won Sick
 CS Nat. Prod. Inst., Seoul Natl. Univ., Seoul, 110-460, S. Korea
 SO Han'guk Saenghwa Hakhoechi (1990), 23(1), 122-7
 CODEN: KBCJAK; ISSN: 0368-4881
 DT Journal
 LA English

L71 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2003 ACS
 AN 1989:225383 CAPLUS
 DN 110:225383
 TI Methyl eugenol: laboratory evaluation in animals
 AU Barbosa, P. P. P.; Teixeira, J. R. M.; Melo, M. F. B.; Gusmao, Q. M. W. B.
 CS Esc. Cienc. Med., Univ. Fed. Alagoas, Alagoas, Brazil
 SO Revista Brasileira de Anestesiologia (1988), 38(6), 393-7
 CODEN: RBANAV; ISSN: 0034-7094
 DT Journal
 LA Portuguese

L71 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2003 ACS
 AN 1982:504098 CAPLUS
 DN 97:104098
 TI The pharmacological effects of a ligroin extract of nutmeg (*Myristica fragrans*)

AU Sherry, C. J.; Ray, L. E.; Herron, R. E.
CS Dep. Biol., Texas A and M Univ., College Station, TX, 77843, USA
SO Journal of Ethnopharmacology (1982), 6(1), 61-6
CODEN: JOETD7; ISSN: 0378-8741
DT Journal
LA English

L71 ANSWER 7 OF 8 CAPLUS COPYRIGHT 2003 ACS
AN 1982:192951 CAPLUS
DN 96:192951
TI Pharmacological studies on methyleugenol
AU Jiang, Ying; Liu, Guoqing; Ma, Junru; Xie, Lin; Wu, Huiqiu
CS Dep. Pharmacol., Nanjing Coll. Pharm., Nanjing, Peop. Rep. China
SO Yaoxue Xuebao (1982), 17(2), 87-92
CODEN: YHHPAL; ISSN: 0513-4870
DT Journal
LA Chinese

L71 ANSWER 8 OF 8 CAPLUS COPYRIGHT 2003 ACS
AN 1973:11566 CAPLUS
DN 78:11566
TI Pharmacologic evaluation of 3,4-dimethoxyphenylpropenes and
3,4-dimethoxyphenylpropanediols
AU Engelbrecht, J. A.; Long, J. P.; Nichols, D. E.; Barfknecht, C. F.
CS Coll. Med., Univ. Iowa, Iowa City, IA, USA
SO Archives Internationales de Pharmacodynamie et de Therapie (1972), 199(2),
226-44
CODEN: AIPTAK; ISSN: 0003-9780
DT Journal
LA English

=> d 171 3 5 6 all

L71 ANSWER 3 OF 8 CAPLUS COPYRIGHT 2003 ACS
AN 1996:287269 CAPLUS
DN 125:1102
TI Synthesis and pharmacological activity of a eugenol derivative
AU Costa, J. A.; Oliveira, R. A. G.; Barbosa, J. M.; Souza Brito, A. R. M.
CS Lab. Tecnologia Farmaceutica, UFPB, Joao Pessoa, Brazil
SO Revista Brasileira de Farmacia (1994), 75(2), 40-5
CODEN: RBFAAH; ISSN: 0370-372X
PB Associacao Brasileira de Farmaceuticos
DT Journal
LA Portuguese
CC 1-11 (Pharmacology)
Section cross-reference(s): 26
AB The aim of this work was the synthesis of a natural pharmacol. active
substance. The target compd. could be prep'd. by an oxidative coupling
reaction involving a starting material also found in nature. Eugenol, an
allyl phenol widely used as a dental local anesthetic, was obtained by a
soxhlet extn. of cloves oil from Caryophyllus aromaticus. Eugenol, prep'd.
by purifn. of the crude oil, was dimerized using potassium ferricyanide,
giving dehydrodieugenol (DDE), a substance previously isolated from
plants. The two phenolic groups were methylated with di-Me sulfate giving
di-O-methyldehydrodieugenol (DMDDE). Pharmacol. evaluation of DMDDDE in
mice showed that it has a CNS-depressant effect, characterized by general
sluggishness of the animal. It potentiated the **sleep** induced by
sodium pentobarbital (which confirms its depressant activity) and also
presented an analgesic effect after chem., mech. and thermal nociceptives
stimulus. Furthermore, 50% of the exptl. animals were protected against
pentylentetrazol-induced convulsion and survived. These data confirmed the

ST central depressant activity of DMDDE.
ST eugenol deriv prepn central nervous depressant; dimethyldehydrodieugenol
prepn central nervous depressant
IT Analgesics
Anticonvulsants and Antiepileptics
Nervous system depressants
Sleep
(dimethyldehydrodieugenol prepn. and pharmacol. activity)
IT 13417-56-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation)
(dimethyldehydrodieugenol prepn. and pharmacol. activity)
IT 97-53-0, Eugenol
RL: RCT (Reactant); RACT (Reactant or reagent)
(dimethyldehydrodieugenol prepn. and pharmacol. activity)
IT 4433-08-3P, Dehydrodieugenol
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(dimethyldehydrodieugenol prepn. and pharmacol. activity)

L71 ANSWER 5 OF 8 CAPLUS COPYRIGHT 2003 ACS
AN 1989:225383 CAPLUS
DN 110:225383
TI Methyl eugenol: laboratory evaluation in animals
AU Barbosa, P. P. P.; Teixeira, J. R. M.; Melo, M. F. B.; Gusmao, Q. M. W. B.
CS Esc. Cienc. Med., Univ. Fed. Alagoas, Alagoas, Brazil
SO Revista Brasileira de Anestesiologia (1988), 38(6), 393-7
CODEN: RBANAV; ISSN: 0034-7094
DT Journal
LA Portuguese
CC 1-11 (Pharmacology)
AB Me Eugenol, an essential oil fraction obtained from *Caryophyllum
aromaticus*, caused central depressing effects with significant hypnotic
and myorelaxing action in doses of 40 mg/kg, i.p., for rats and 5 mg/kg,
i.v., for rabbits and dogs, rapid induction and satisfactory duration of
sleep (118.4 s and 47.3 min resp.) in rats, and **sleep**
time between 9-12 min in dogs. Anesthetic evolution in dogs was
satisfactory, followed by rapid recovery and movement. Me eugenol (20
.mu.g/mL) reduced the atrial muscular cardiac force of contraction (50%)
in 20 min. The addn. of Me eugenol to isolated rat diaphragm-nerve
preps. produced muscular contraction blockade under direct and indirect
stimulation.
ST methyl eugenol hypnotic muscle relaxant
IT Anesthetics
Hypnotics and Sedatives
Muscle relaxants
(Me eugenol)
IT 93-15-2, Methyl eugenol
RL: BIOL (Biological study)
(hypnotic and muscle-relaxant activities of)

L71 ANSWER 6 OF 8 CAPLUS COPYRIGHT 2003 ACS
AN 1982:504098 CAPLUS
DN 97:104098
TI The pharmacological effects of a ligroin extract of nutmeg (*Myristica
fragrans*)
AU Sherry, C. J.; Ray, L. E.; Herron, R. E.
CS Dep. Biol., Texas A and M Univ., College Station, TX, 77843, USA
SO Journal of Ethnopharmacology (1982), 6(1), 61-6
CODEN: JOETD7; ISSN: 0378-8741
DT Journal

LA English
CC 1-11 (Pharmacology)
Section cross-reference(s): 11, 63
AB A ligroin ext. of nutmeg (*Myristica fragrans*) increased the duration of light and deep **sleep** in the young chicken. The presence of trimyristin [555-45-3] tended to increase the effect of the ext. The ext. did not contain detectable amts. of myristicin [607-91-0], elemicin [487-11-6], safrole [94-59-7], or eugenol [97-53-0], which either individually or collectively have been suggested to be the active agents of nutmeg.
ST nutmeg ext pharmacol; psychotropic nutmeg ext
IT Myristica
 (ext. of, compn. and pharmacol. of)
IT Psychotropics
 (nutmeg ext.)
IT 94-59-7 97-53-0 487-11-6 607-91-0
RL: BIOL (Biological study)
 (nutmeg psychotropic activity in relation to)
IT 555-45-3
RL: BIOL (Biological study)
 (nutmeg psychotropic activity potentiation by)

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

L1 43 S CEDROL
L2 33 S PATCHOULI
L3 36 S SANTALOL
L4 31 S BISABOLOL
 E BISABOLOL
L5 2 S VETIVEROL
L6 30 S SCLAREOL
L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

 E SLEEP
L10 15616 S E3
 E SEDATIVE
L11 12852 S E3-E9
 E RELAXATION
L12 217668 S E3 OR E8
 E NARCOTIC
L13 10398 S E3 OR E12
 E HYPNOTIC
L14 9427 S E3 OR E10
 E SOMULENCE
 E SOMUL
 E SOMNIA
 E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3
L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8

L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
L27 1 S 38419-75-9/RN
SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

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L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16
L34 0 S L17 AND L10
L35 0 S L17 AND L11
L36 0 S L17 AND L13
L37 0 S L17 AND L15
L38 0 S L18 AND L10
L39 0 S L18 AND L13
L40 0 S L19 AND L10
L41 0 S LL18 AND L12
L42 0 S L18 AND L12
L43 1 S L19 AND L12
L44 1 S 43 1 ALL
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L46 0 S L20 AND L10
L47 0 S L20 AND L12
L48 1 S L21 AND L10
L49 0 S L22 AND L12
L50 0 S L25 AND L10
E NERVOUS
E NERVOUS
L51 164186 S E3-E7
L52 1 S L51 AND L16

FILE 'REGISTRY' ENTERED AT 16:16:22 ON 15 JUL 2003
E FARNESOL
L53 75 S E3
E EUGENOL
L54 165 S E3
L55 4 S GERANYL LINALOOL
E CEDRENOL
L56 9 S E3
E ISOPYTOL
E ISOPHYTOL
L57 6 S E3
E NEROLIDOL
L58 41 S E3

FILE 'CPLUS' ENTERED AT 16:21:24 ON 15 JUL 2003
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L60 0 S L56 AND L12
L61 0 S L56 AND L15
L62 90 S L56
L63 1 S L62 AND L10
L64 0 S L62 AND L12
L65 3275 S L53
L66 1 S L65 AND L10

L67 9 S L65 AND L12
L68 0 S L65 AND L13
L69 0 S L65 AND L15
L70 8961 S L54
L71 8 S L70 AND L10

=> s 170 and l12
L72 33 L70 AND L12

=> d 172 not 171
L71 IS NOT VALID HERE
For an explanation, enter "HELP DISPLAY".

=> s 172 not 171
L73 33 L72 NOT L71

=> s 172 10-33
MISSING OPERATOR L72 10-33
The search profile that was entered contains terms or
nested terms that are not separated by a logical operator.

=> d 172 10-33

L72 ANSWER 10 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1997:531315 CAPLUS
DN 127:242982
TI Eugenolol: a eugenol-derived .beta.-adrenoceptor blocker with partial
.beta.2-agonist and calcium mobilization inhibition associated
vasorelaxant activities
AU Chen, Sheue-Jiun; Huang, Yeun-Chih; Wu, Bin-Nan; Chen, Ing-Jun
CS Department of Pharmacology, Kaohsiung Medical College, Kaohsiung, 807,
Taiwan
SO Drug Development Research (1997), 40(3), 239-250
CODEN: DDREDK; ISSN: 0272-4391
PB Wiley-Liss
DT Journal
LA English

L72 ANSWER 11 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1996:350392 CAPLUS
DN 125:47250
TI Electron Spin Resonance Studies of Reorientational Motion in Glass-Forming
Liquids
AU Kowert, Bruce A.; Higgins, Edward J.; Mariencheck, William I.; Stemmler,
Timothy L.; Kantorovich, Vladimir
CS Department of Chemistry, Saint Louis University, Saint Louis, MO, 63103,
USA
SO Journal of Physical Chemistry (1996), 100(27), 11211-11217
CODEN: JPCHAX; ISSN: 0022-3654
PB American Chemical Society
DT Journal
LA English

L72 ANSWER 12 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1996:193913 CAPLUS
DN 124:275865
TI Experimental study of dielectric **relaxation** in supercooled
alcohols and polyols
AU Murthy, S. S. N.
CS Sch. Phys. Sci., Jawaharlal Nehru Univ., New Delhi, 110067, India
SO Molecular Physics (1996), 87(3), 691-709
CODEN: MOPHAM; ISSN: 0026-8976

PB Taylor & Francis
DT Journal
LA English

L72 ANSWER 13 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1995:990814 CAPLUS
DN 124:9689
TI Extrusion-moldable polyolefin resins suitable for moldings having complicated profiles
IN Tsuruoka, Masayuki; Nakagawa, Susumu; Hirano, Koki
PA Idemitsu Petrochemical Co, Japan
SO Jpn. Kokai Tokkyo Koho, 14 pp.
CODEN: JKXXAF

DT Patent
LA Japanese

FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|--------------------|------|----------|-----------------|----------|
| ----- | --- | ----- | ----- | ----- |
| PI JP 07247318 | A2 | 19950926 | JP 1994-38168 | 19940309 |
| PRAI JP 1994-38168 | | 19940309 | | |

L72 ANSWER 14 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1993:410647 CAPLUS
DN 119:10647
TI Phosphorus-31 NMR spectroscopy in wood chemistry. Part IV. Lignin models: spin lattice **relaxation** times and solvent effects in phosphorus-31 NMR
AU Argyropoulos, Dimitris S.; Bolker, Henry I.; Heitner, Cyril; Archipov, Yuri
CS Dep. Chem., McGill Univ., Montreal, QC, H3A 2A7, Can.
SO Holzforschung (1993), 47(1), 50-6
CODEN: HOLZAZ; ISSN: 0018-3830
DT Journal
LA English

L72 ANSWER 15 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1988:556844 CAPLUS
DN 109:156844
TI Structural **relaxation** mechanisms in liquid eugenol. A depolarized light scattering study
AU Bezot, P.; Hesse-Bezot, C.; Roynard, D.; Jeanneaux, F.
CS Lab. Phys. Matiere Condens., Nice, 06034, Fr.
SO Journal of Chemical Physics (1988), 89(1), 1-5
CODEN: JCPSA6; ISSN: 0021-9606
DT Journal
LA English

L72 ANSWER 16 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1987:648750 CAPLUS
DN 107:248750
TI Proton longitudinal **relaxation** times of carbon-13 isotopomers
AU Bigler, Peter
CS Inst. Org. Chem., Univ. Berne, Bern, 3012, Switz.
SO Journal of Magnetic Resonance (1969-1992) (1987), 75(1), 162-6
CODEN: JOMRA4; ISSN: 0022-2364
DT Journal
LA English

L72 ANSWER 17 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1985:143009 CAPLUS
DN 102:143009
TI Relaxant effects on tracheal and ileal smooth muscles of the guinea pig

AU Reiter, M.; Brandt, W.
CS Inst. Pharmakol. Toxikol., Tech. Univ. Muenchen, Munich, D-8000/40, Fed.
Rep. Ger.
SO Arzneimittel-Forschung (1985), 35(1A), 408-14
CODEN: ARZNAD; ISSN: 0004-4172
DT Journal
LA English

L72 ANSWER 18 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1983:510521 CAPLUS
DN 99:110521
TI Quantitative analysis of clove oil by NMR spectrometry
AU Chiang, Hung Cheh; Wang, Pei Lein; Huang, Keh Feng
CS Inst. Chem., Natl. Taiwan Norm. Univ., Taipei, 117, Taiwan
SO Journal of the Chinese Chemical Society (Taipei, Taiwan) (1983), 30(2),
117-20
CODEN: JCCTAC; ISSN: 0009-4536
DT Journal
LA English

L72 ANSWER 19 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1983:400161 CAPLUS
DN 99:161
TI Chemostructural requirement for centrally acting muscle relaxant effect of
magnolol and honokiol, neolignane derivatives
AU Watanabe, Hiroshi; Watanabe, Kazuo; Hagino, Koji
CS Res. Inst. Wakan-yaku, Toyama Med. Pharm. Univ., Toyama, 930-01, Japan
SO Journal of Pharmacobio-Dynamics (1983), 6(3), 184-90
CODEN: JOPHDQ; ISSN: 0386-846X
DT Journal
LA English

L72 ANSWER 20 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1982:515659 CAPLUS
DN 97:115659
TI Acoustic and viscoelastic **relaxation** in liquid eugenol
AU Karabaev, M. K.; Turdyev, N. Sh.
CS Otd. Teplofiz., Tashkent, USSR
SO Izvestiya Akademii Nauk UzSSR, Seriya Fiziko-Matematicheskikh Nauk (1982),
(2), 50-1
CODEN: IUZFAU; ISSN: 0131-8012
DT Journal
LA Russian

L72 ANSWER 21 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1978:624673 CAPLUS
DN 89:224673
TI Dielectric **relaxation** in dilute solutions of some hydroxy
compounds
AU Hanna, Faika Fahmy; Bishai, Augenie Michael
CS Arab Dev. Inst., Tripoli, Libya
SO Zeitschrift fuer Physikalische Chemie (Leipzig) (1978), 259(5), 849-55
CODEN: ZPCLAH; ISSN: 0372-9680
DT Journal
LA English

L72 ANSWER 22 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1976:502884 CAPLUS
DN 85:102884
TI The effects of temperature and pressure on the complex dielectric
permittivity of liquid eugenol and glycerol
AU Scaife, W. G. S.

CS Eng. Sch., Trinity Coll., Dublin, Ire.
SO Journal of Physics D: Applied Physics (1976), 9(10), 1489-99
CODEN: JPAPBE; ISSN: 0022-3727
DT Journal
LA English

L72 ANSWER 23 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1976:427707 CAPLUS
DN 85:27707
TI Dielectric **relaxation** in eugenol
AU Alper, Turhan; Barlow, A. John; Kim, Min G.
CS Dep. Electron. Electr. Eng., Univ. Glasgow, Glasgow, UK
SO Journal of the Chemical Society, Faraday Transactions 2: Molecular and Chemical Physics (1976), 72(5), 934-40
CODEN: JCFTBS; ISSN: 0300-9238
DT Journal
LA English

L72 ANSWER 24 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1975:154989 CAPLUS
DN 82:154989
TI Viscoelastic **relaxation** in supercooled eugenol
AU Kim, Min Gon
CS Dep. Electron. Electr. Eng., Univ. Glasgow, Glasgow, UK
SO Journal of the Chemical Society, Faraday Transactions 2: Molecular and Chemical Physics (1975), 71(3), 415-22
CODEN: JCFTBS; ISSN: 0300-9238
DT Journal
LA English

L72 ANSWER 25 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1974:137424 CAPLUS
DN 80:137424
TI Compressional study of alcohols through pseudo-Grueneisen parameter
AU Tandon, Uma S.
CS Dep. Phys., Univ. Allahabad, Allahabad, India
SO Proc. Nucl. Phys. Solid State Phys. Symp., 17th (1973), Meeting Date 1972, Volume C, 309-12 Publisher: Phys. Comm. Dep. At. Energy, Bombay, India.
CODEN: 27GNAE
DT Conference
LA English

L72 ANSWER 26 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1974:71472 CAPLUS
DN 80:71472
TI Changes of viscoelastic properties of poly(methyl methacrylate) soaked in various organic solvents
AU Yanaru, Ritsuo
CS Kyushu Dent. Coll., Kitakyushu, Japan
SO Kyushu Shika Gakkai Zasshi (1973), 26(5), 224-51
CODEN: KSGZA3; ISSN: 0368-6833
DT Journal
LA Japanese

L72 ANSWER 27 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1974:49471 CAPLUS
DN 80:49471
TI Dielectric properties of lignin
AU Norimoto, Misato; Nakatsubo, Fumiaki; Yamada, Tadashi
CS Wood Res. Inst., Kyoto Univ., Kyoto, Japan
SO Zairyo (1973), 22(241), 937-42

DT CODEN: ZARYAQ; ISSN: 0514-5163
LA Journal
LA Japanese

L72 ANSWER 28 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1973:529186 CAPLUS
DN 79:129186
TI Pressure dependence of ultrasonic absorption in eugenol and carbon tetrachloride
AU Kor, S. K.; Pandey, S. K.
CS Dep. Phys., Univ. Allahabad, Allahabad, India
SO Journal of the Physical Society of Japan (1973), 35(4), 1175-8
CODEN: JUPSAU; ISSN: 0031-9015
DT Journal
LA English

L72 ANSWER 29 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1969:81945 CAPLUS
DN 70:81945
TI Effect of pressure on the complex permittivity of eugenol
AU Scaife, W. G.
CS Trinity Coll., Dublin, Ire.
SO National Academy of Sciences-National Research Council, Publication (1968), No. 1578, 70-80
CODEN: NASRAE; ISSN: 0547-8464
DT Journal
LA English

L72 ANSWER 30 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1964:12432 CAPLUS
DN 60:12432
OREF 60:2227a-d
TI Neuromuscular blocking action of a general anesthetic, the N,N-diethylamide of 2-methoxy-4-allylphenoxyacetic acid (Estil)
AU Malafaya-Baptista, A.; Guimaraes, S.; Rodrigues-Pereira, E.
CS Univ. Oporto, Port.
SO Archives Internationales de Pharmacodynamie et de Therapie (1963), 145(1-2), 44-50
CODEN: AIPTAK; ISSN: 0003-9780
DT Journal
LA English

L72 ANSWER 31 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1962:472361 CAPLUS
DN 57:72361
OREF 57:14403f-h
TI A new anesthetic, 2-methoxyl-4-allylphenoxyacetic acid diethylamide (Detrovel) in the clearing and curetting of the uterus following abortions
AU Neumann, E.; Bernadin, D.; Couturier, J. C.
CS Clin. Obstet., Lyons, Fr.
SO Bull. Federation Soc. Gynecol. Obstet. Langue Franc. (1962), 14, 147-8
DT Journal
LA Unavailable

L72 ANSWER 32 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1962:57028 CAPLUS
DN 56:57028
OREF 56:10875f-h
TI Gas analyses in umbilical cord blood after obstetric procedures
AU Kittel, E.; Mueller-Plathe, O.; Schmolling, E.
CS Univ.-Frauenklinik, Hamburg-Eppendorf, Germany
SO Klinische Wochenschrift (1961), 39, 911-13

CODEN: KIWOAZ; ISSN: 0023-2173
DT Journal
LA Unavailable

L72 ANSWER 33 OF 33 CAPLUS COPYRIGHT 2003 ACS
AN 1954:70272 CAPLUS
DN 48:70272
OREF 48:12488i,12489a-b
TI Dielectric **relaxation** time and association. III. The role of intramolecular and intermolecular hydrogen-bond formation of ortho-substituted phenols in relation to their concentration in nonpolar carbon tetrachloride
AU Fischer, Erich
CS Univ. Ankara, Turk.
SO Zeitschrift fuer Naturforschung (1954), 9a, 360-5
CODEN: ZNTFA2; ISSN: 0372-9516
DT Journal
LA Unavailable

=> s 155
L74 123 L55

=> s 174 and 110
L75 1 L74 AND L10

=> d 175

L75 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1992:171087 CAPLUS
DN 116:171087
TI Effects of olfactory stimulation with jasmin and its component chemicals on the duration of pentobarbital-induced **sleep** in mice
AU Tsuchiya, T.; Tanida, M.; Uenoyama, S.; Nakayama, Y.
CS Shiseido Res. Cent., Yokohama, 223, Japan
SO Life Sciences (1992), 50(15), 1097-102
CODEN: LIIFSAK; ISSN: 0024-3205
DT Journal
LA English

=> d 175 1 all

L75 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1992:171087 CAPLUS
DN 116:171087
TI Effects of olfactory stimulation with jasmin and its component chemicals on the duration of pentobarbital-induced **sleep** in mice
AU Tsuchiya, T.; Tanida, M.; Uenoyama, S.; Nakayama, Y.
CS Shiseido Res. Cent., Yokohama, 223, Japan
SO Life Sciences (1992), 50(15), 1097-102
CODEN: LIIFSAK; ISSN: 0024-3205
DT Journal
LA English
CC 13-6 (Mammalian Biochemistry)
Section cross-reference(s): 62
AB The effect of olfactory stimulation with jasmin and its component chems. on pentobarbital **sleep** time was investigated using mice in order to det. which component of jasmin influences pentobarbital **sleep** time via olfactory stimulation. **Sleep** time was defined as the time elapsed between i.p. pentobarbital administration and the first time that the animal was able to spontaneously right itself. **Sleep**

time was significantly decreased by olfactory stimulation with jasmin, and also by one of the fractions obtained by fractional distn. at 150 degree.C and 0.1 mmHg. The fraction which influenced the **sleep** time was found to consist of benzyl benzoate, isophytol, geranyl linalool, phytol and phytol acetate, which were identified using gas chromatog. with mass and IR spectrometry. In expts. using authentic samples of these components, phytol significantly shortened the pentobarbital **sleep** time, while the others had no effect. Phytol is the component of jasmin which reduces the duration of pentobarbital-induced **sleep**.

ST **sleep** pentobarbital jasmin phytol drug interaction; olfactory system **sleep** pentobarbital jasmin phytol

IT **sleep**
(jasmin inhibition of pentobarbital-induced, olfactory stimulation in)

IT Essential oils
RL: BIOL (Biological study)
(jasmine, Jasminum grandiflorum abs., pentobarbital-induced **sleep** inhibition by, olfactory stimulation in)

IT Nervous system
(olfactory system, jasmin stimulation of, pentobarbital-induced **sleep** inhibition by)

IT 76-74-4, Pentobarbital
RL: BIOL (Biological study)
(jasmin inhibition of **sleep** stimulation by, olfactory stimulation in)

IT 120-51-4, Benzyl benzoate 140-11-4, Benzyl acetate 150-86-7, Phytol 505-32-8, Isophytol 1113-21-9, Geranyl linalool 10236-16-5, Phytol acetate
RL: BIOL (Biological study)
(pentobarbital **sleep** time response to, as jasmin component, olfactory stimulation in relation to)

=> s 157
L76 380 L57

=> s 176 and 110
L77 1 L76 AND L10

=> d 177

L77 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1992:171087 CAPLUS
DN 116:171087
TI Effects of olfactory stimulation with jasmin and its component chemicals on the duration of pentobarbital-induced **sleep** in mice
AU Tsuchiya, T.; Tanida, M.; Uenoyama, S.; Nakayama, Y.
CS Shiseido Res. Cent., Yokohama, 223, Japan
SO Life Sciences (1992), 50(15), 1097-102
CODEN: LIFSAK; ISSN: 0024-3205
DT Journal
LA English

=> s 176 and 115
L78 0 L76 AND L15

=> s 158
L79 2587 L58

=> s 179 and 110
L80 2 L79 AND L10

=> d 180 1-2

L80 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS
AN 2001:753166 CAPLUS
DN 135:308609
TI Perfume compositions for memory improvement
IN Tanisawa, Shigeji; Suga, Chihoko; Goto, Masahiro; Okuda, Takehiro
PA Pola Chemical Industries, Inc., Japan
SO Jpn. Kokai Tokkyo Koho, 5 pp.
CODEN: JKXXAF
DT Patent
LA Japanese
FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|----------------|-------|----------|-----------------|----------|
| | ----- | ----- | ----- | ----- | ----- |
| PI | JP 2001288493 | A2 | 20011016 | JP 2000-103001 | 20000405 |
| PRAI | JP 2000-103001 | | 20000405 | | |

L80 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS
AN 2001:124383 CAPLUS
DN 134:183320
TI Perfumes and their compositions for stress alleviation
IN Tanisawa, Shigeji; Suga, Chihoko; Goto, Masahiro; Okuda, Takehiro;
Ishitoya, Toyomasa
PA Pola Chemical Industries, Inc., Japan
SO Jpn. Kokai Tokkyo Koho, 5 pp.
CODEN: JKXXAF
DT Patent
LA Japanese
FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|----------------|-------|----------|-----------------|----------|
| | ----- | ----- | ----- | ----- | ----- |
| PI | JP 2001049286 | A2 | 20010220 | JP 1999-221887 | 19990805 |
| PRAI | JP 1999-221887 | | 19990805 | | |

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(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

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|----|----|-------------|
| L1 | 43 | S CEDROL |
| L2 | 33 | S PATCHOULI |
| L3 | 36 | S SANTALOL |
| L4 | 31 | S BISABOLOL |
| | E | BISABOLOL |
| L5 | 2 | S VETIVEROL |
| L6 | 30 | S SCLAREOL |
| L7 | 0 | S GLOBUOL |
| L8 | 9 | S GLOBULOL |
| L9 | 11 | S GUAIOL |

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

| | | |
|-----|--------------|-------------|
| | E SLEEP | |
| L10 | 15616 | S E3 |
| | E SEDATIVE | |
| L11 | 12852 | S E3-E9 |
| | E RELAXATION | |
| L12 | 217668 | S E3 OR E8 |
| | E NARCOTIC | |
| L13 | 10398 | S E3 OR E12 |
| | E HYPNOTIC | |

L14 9427 S E3 OR E10
 E SOMULENCE
 E SOMUL
 E SOMNIA
 E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3
L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8
L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
L27 1 S 38419-75-9/RN
 SET NOTICE 1 DISPLAY
 SET NOTICE LOGIN DISPLAY

FILE 'CAPPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003
L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16
L34 0 S L17 AND L10
L35 0 S L17 AND L11
L36 0 S L17 AND L13
L37 0 S L17 AND L15
L38 0 S L18 AND L10
L39 0 S L18 AND L13
L40 0 S L19 AND L10
L41 0 S LL18 AND L12
L42 0 S L18 AND L12
L43 1 S L19 AND L12
L44 1 S 43 1 ALL
L45 0 S L19 AND L15
L46 0 S L20 AND L10
L47 0 S L20 AND L12
L48 1 S L21 AND L10
L49 0 S L22 AND L12
L50 0 S L25 AND L10
 E NERVIOS
 E NERVOUS
L51 164186 S E3-E7
L52 1 S L51 AND L16

FILE 'REGISTRY' ENTERED AT 16:16:22 ON 15 JUL 2003
 E FARNESOL
L53 75 S E3
 E EUGENOL
L54 165 S E3
L55 4 S GERANYL LINALOOL
 E CEDRENOL
L56 9 S E3
 E ISOPYTOL
 E ISOPHYTOL

L57 6 S E3
 E NEROLIDOL
L58 41 S E3

FILE 'CAPLUS' ENTERED AT 16:21:24 ON 15 JUL 2003

L59 1 S L56 AND L10
L60 0 S L56 AND L12
L61 0 S L56 AND L15
L62 90 S L56
L63 1 S L62 AND L10
L64 0 S L62 AND L12
L65 3275 S L53
L66 1 S L65 AND L10
L67 9 S L65 AND L12
L68 0 S L65 AND L13
L69 0 S L65 AND L15
L70 8961 S L54
L71 8 S L70 AND L10
L72 33 S L70 AND L12
L73 33 S L72 NOT L71
L74 123 S L55
L75 1 S L74 AND L10
L76 380 S L57
L77 1 S L76 AND L10
L78 0 S L76 AND L15
L79 2587 S L58
L80 2 S L79 AND L10

=> d 148

L48 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian

=> d 148 all

L48 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian
CC 1-11 (Pharmacology)
AB In expts. on male mice and rats some behavioral effects of the diterpene sclareol glucol (SG) were studied. SG in higher doses (50, 100 mg/kg) enhanced the sensitivity to touch and pain, shortened the duration of hexobarbital sleep, stimulated exploratory behavior in rats in open field, exerted no anti-convulsive effect and increased the mortality from pentylenetetrazole seizures. In higher doses (300, 500, and 1000 mg/kg) SG induced seizures.

ST sclareol glucol behavior nervous system
IT Behavior
Nervous system
(sclareol glucol effect on)
IT 38419-75-9
RL: PRP (Properties)
(behavioral and nervous systems effects of)

=> e stress
E1 1 STRESP/BI
E2 1 STRESPTOMYCES/BI
E3 393704 --> STRESS/BI
E4 2 STRESS1/BI
E5 1 STRESS2/BI
E6 1 STRESS3/BI
E7 1 STRESS4/BI
E8 7 STRESS70/BI
E9 1 STRESS70C/BI
E10 1 STRESS7WAS/BI
E11 5 STRESSA/BI
E12 8 STRESSABILITY/BI

=> s e3
L81 393704 STRESS/BI

=> s l81 and l16
L82 5 L81 AND L16

=> d l82 1-5

L82 ANSWER 1 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN 2003:146475 CAPLUS
DN 138:192839
TI Stress-relieving hair-styling preparations containing sesquiterpene alcohols
IN Nagashima, Yoshinao; Yata, Sachihiro
PA Kao Corp., Japan
SO Jpn. Kokai Tokkyo Koho, 9 pp.
CODEN: JKXXAF

DT Patent
LA Japanese

FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------------|------|----------|-----------------|----------|
| PI JP 2003055161 | A2 | 20030226 | JP 2001-244909 | 20010810 |
| PRAI JP 2001-244909 | | 20010810 | | |

L82 ANSWER 2 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN 2001:597788 CAPLUS
DN 135:170507
TI Autonomic-controlling agents containing sesquiterpene alcohols
IN Nagashima, Yoshinao; Sugata, Keiichi; Yada, Yukihiro; Fukuda, Kazuyuki
PA Kao Corp., Japan
SO PCT Int. Appl., 48 pp.
CODEN: PIXXD2

DT Patent
LA Japanese

FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------|------|----------|-----------------|----------|
| PI WO 2001058435 | A1 | 20010816 | WO 2001-JP928 | 20010209 |

W: JP, US
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
PT, SE, TR
EP 1170005 A1 20020109 EP 2001-902822 20010209
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, FI
US 2002151600 A1 20021017 US 2001-972887 20011010
PRAI JP 2000-38260 A 20000210
WO 2001-JP928 W 20010209

RE.CNT 12 THERE ARE 12 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN 2000:35577 CAPLUS
DN 132:332157
TI Physiological effects of volatile components in forest
AU Sawada, Kazuhiko; Komaki, Ryoichi; Yamashita, Yoshikuni; Suzuki, Yasushi;
Ishiyama, Seiichi
CS Cosmetic Lab., Kanebo Co. Ltd., Japan
SO Nippon Aji to Nioi Gakkaishi (1999), 6(3), 465-468
CODEN: NNGAEW; ISSN: 1340-4806
PB Nippon Aji to Nioi Gakkai
DT Journal
LA Japanese

L82 ANSWER 4 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN 1999:493051 CAPLUS
DN 131:348943
TI Antifungal activity to Phytophthora infestans of sesquiterpenoids from
infected potato tubers
AU Engstrom, K.; Widmark, A. K.; Brishammar, S.; Helmersson, S.
CS Department of Chemistry, Swedish University of Agricultural Sciences,
Uppsala, S-750, Swed.
SO Potato Research (1999), 42(1), 43-50
CODEN: PORHBW; ISSN: 0014-3065
PB European Association for Potato Research
DT Journal
LA English

RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L82 ANSWER 5 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN 1994:187222 CAPLUS
DN 120:187222
TI The effect of a long-term water **stress** on the metabolism and
emission of terpenes of the foliage of Cupressus sempervirens
AU Yani, A.; Pauly, G.; Faye, M.; Salin, F.; Gleizes, M.
CS Lab. Physiol. Cell. Veg., Univ. Bordeaux I, Talence, 33405, Fr.
SO Plant, Cell and Environment (1993), 16(8), 975-81
CODEN: PLCEDV; ISSN: 0140-7791
DT Journal
LA English

=> d 182 3 all

L82 ANSWER 3 OF 5 CAPLUS COPYRIGHT 2003 ACS
AN 2000:35577 CAPLUS
DN 132:332157
TI Physiological effects of volatile components in forest
AU Sawada, Kazuhiko; Komaki, Ryoichi; Yamashita, Yoshikuni; Suzuki, Yasushi;
Ishiyama, Seiichi

CS Cosmetic Lab., Kanebo Co. Ltd., Japan
SO Nippon Aji to Nioi Gakkaishi (1999), 6(3), 465-468
CODEN: NNGAEW; ISSN: 1340-4806
PB Nippon Aji to Nioi Gakkai
DT Journal
LA Japanese
CC 11-8 (Plant Biochemistry)
Section cross-reference(s): 13
AB The physiol. effect such as **stress** relief of hiba forest
volatile components such as monoterpenes is studied.
ST hiba forest volatile component physiol function
IT Forests
 (hiba; physiol. effects of volatile components in forest)
IT Thujopsis dolabrata
Volatile substances
 (physiol. effects of volatile components in forest)
IT Monoterpene
RL: BSU (Biological study, unclassified); GOC (Geological or astronomical
occurrence); BIOL (Biological study); OCCU (Occurrence)
 (physiol. effects of volatile components in forest)
IT 77-53-2, Cedrol 80-56-8, .alpha.-Pinene 13466-78-9,
.DELTA.3-Carene
RL: BSU (Biological study, unclassified); GOC (Geological or astronomical
occurrence); BIOL (Biological study); OCCU (Occurrence)
 (physiol. effects of volatile components in forest)

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

L1 43 S CEDROL
L2 33 S PATCHOULI
L3 36 S SANTALOL
L4 31 S BISABOLOL
 E BISABOLOL
L5 2 S VETIVEROL
L6 30 S SCLAREOL
L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

 E SLEEP
L10 15616 S E3
 E SEDATIVE
L11 12852 S E3-E9
 E RELAXATION
L12 217668 S E3 OR E8
 E NARCOTIC
L13 10398 S E3 OR E12
 E HYPNOTIC
L14 9427 S E3 OR E10
 E SOMULENCE
 E SOMUL
 E SOMNIA
 E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3

L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8
L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003
L27 1 S 38419-75-9/RN
 SET NOTICE 1 DISPLAY
 SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003
L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16
L34 0 S L17 AND L10
L35 0 S L17 AND L11
L36 0 S L17 AND L13
L37 0 S L17 AND L15
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L40 0 S L19 AND L10
L41 0 S L18 AND L12
L42 0 S L18 AND L12
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L47 0 S L20 AND L12
L48 1 S L21 AND L10
L49 0 S L22 AND L12
L50 0 S L25 AND L10
 E NERVOUS
 E NERVOUS
L51 164186 S E3-E7
L52 1 S L51 AND L16

FILE 'REGISTRY' ENTERED AT 16:16:22 ON 15 JUL 2003
E FARNESOL
L53 75 S E3
 E EUGENOL
L54 165 S E3
L55 4 S GERANYL LINALOOL
 E CEDRENOL
L56 9 S E3
 E ISOPYTOL
 E ISOPHYTOL
L57 6 S E3
 E NEROLIDOL
L58 41 S E3

FILE 'CAPLUS' ENTERED AT 16:21:24 ON 15 JUL 2003
L59 1 S L56 AND L10
L60 0 S L56 AND L12
L61 0 S L56 AND L15
L62 90 S L56

L63 1 S L62 AND L10
L64 0 S L62 AND L12
L65 3275 S L53
L66 1 S L65 AND L10
L67 9 S L65 AND L12
L68 0 S L65 AND L13
L69 0 S L65 AND L15
L70 8961 S L54
L71 8 S L70 AND L10
L72 33 S L70 AND L12
L73 33 S L72 NOT L71
L74 123 S L55
L75 1 S L74 AND L10
L76 380 S L57
L77 1 S L76 AND L10
L78 0 S L76 AND L15
L79 2587 S L58
L80 2 S L79 AND L10
 E STRESS
L81 393704 S E3
L82 5 S L81 AND L16

=> s l62and l81

MISSING OPERATOR L62AND L81

The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

=> s 162 and 181

L83 0 L62 AND L81

=> s 165 and 181

L84 26 L65 AND L81

=> d 184 10-26

L84 ANSWER 10 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1997:768128 CAPLUS
DN 128:84077
TI Trypanosoma brucei: effects of methoprene and other isoprenoid compounds on procyclic and bloodstream forms in vitro and in mice
AU Harmon, Margaret A.; Scott, Teddy C.; Li, Yuhua; Boehm, Marcus F.; Phillips, Margaret A.; Mangelsdorf, David J.
CS Department of Pharmacology, University of Texas Southwestern Medical Center at Dallas, Dallas, TX, 75235-9041, USA
SO Experimental Parasitology (1997), 87(3), 229-236
CODEN: EXPAAA; ISSN: 0014-4894
PB Academic Press
DT Journal
LA English

L84 ANSWER 11 OF 26 CAPLUS COPYRIGHT 2003 ACS

AN 1997:679260 CAPLUS

DN 128:10189

TI Lovastatin induces apoptosis by inhibiting mitotic and post-mitotic events in cultured mesangial cells
AU Ghosh, Paramita M.; Mott, Glen E.; Ghosh-Choudhury, Nandini; Radnik, Robert A.; Stapleton, Marissa L.; Ghidoni, John J.; Kreisberg, Jeffrey I.
CS Department of Pathology, University of Texas Health Science Center, 7703 Floyd Curl Drive, San Antonio, USA
SO Biochimica et Biophysica Acta (1997), 1359(1), 13-24
CODEN: BBACAO; ISSN: 0006-3002
PB Elsevier

DT Journal
LA English

L84 ANSWER 12 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1996:114474 CAPLUS
DN 124:223355
TI Convergence of three steroid receptor pathways in the mediation of nongenotoxic hepatocarcinogenesis
AU O'Brien, M. L.; Rangwala, S. M.; Henry, K. W.; Weinberger, C.; Crick, D. C.; Waechter, C. J.; Feller, D. R.; Noonan, D. J.
CS Dep. Biochem., University Kentucky, Lexington, KY, 40536, USA
SO Carcinogenesis (1996), 17(2), 185-90
CODEN: CRNGDP; ISSN: 0143-3334
PB Oxford University Press
DT Journal
LA English

L84 ANSWER 13 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1996:40356 CAPLUS
DN 124:79279
TI Is sidestream smoke a stressor?
AU Barbera, Nunziata; Iurato, Maria Pierangela; Geremia, Ernesto; Bernardini, Renato
CS Institutes Pharmacology, University Catania, Catania, I-95125, Italy
SO Indoor Environment (1995), 4(3-4), 157-61
CODEN: IENVEC; ISSN: 1016-4901
PB Karger
DT Journal
LA English

L84 ANSWER 14 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1994:102387 CAPLUS
DN 120:102387
TI Effects of growth regulators on the induction of Crassulacean acid metabolism in the facultative halophyte *Mesembryanthemum crystallinum* L.
AU Dai, Ziyu; Ku, Maurice S. B.; Zhang, Dianzhong; Edwards, Gerald E.
CS Bot. Dep., Washington State Univ., Pullman, WA, 99164-4238, USA
SO Planta (1994), 192(3), 287-94
CODEN: PLANAB; ISSN: 0032-0935
DT Journal
LA English

L84 ANSWER 15 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1992:148339 CAPLUS
DN 116:148339
TI Sorghum isoprenoid pathway responses to manganese concentration
AU Wilkinson, R. E.
CS Dep. Agron., Univ. Georgia, Griffin, GA, 30223-1797, USA
SO Canadian Journal of Plant Science (1991), 71(4), 973-81
CODEN: CPLSAY; ISSN: 0008-4220
DT Journal
LA English

L84 ANSWER 16 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1990:175849 CAPLUS
DN 112:175849
TI Factors influencing the concentration of solanesol in Burley tobacco
AU Burton, H. R.; Leggett, Everett; Phillips, R. E.
CS Dep. Agron., Univ. Kentucky, Lexington, KY, USA
SO Beitrage zur Tabakforschung International (1989), 14(5), 313-20
CODEN: BTAID3; ISSN: 0173-783X
DT Journal

LA English

L84 ANSWER 17 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1990:36226 CAPLUS
DN 112:36226
TI Preparation of antiulcer isoprenoid derivatives and pharmaceutical compositions containing them
IN Yamada, Kazuhiko; Tahara, Yoshiyuki; Toyoda, Masashi; Irino, Osamu; Misaki, Noriyuki
PA Nisshin Flour Milling Co., Ltd., Japan
SO Eur. Pat. Appl., 29 pp.
CODEN: EPXXDW
DT Patent
LA English
FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|---|------|----------|-----------------|----------|
| PI | EP 304842 | A2 | 19890301 | EP 1988-113617 | 19880822 |
| | EP 304842 | A3 | 19910116 | | |
| | EP 304842 | B1 | 19941130 | | |
| | R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE | | | | |
| | JP 02042030 | A2 | 19900213 | JP 1988-206465 | 19880822 |
| | US 4906669 | A | 19900306 | US 1988-234895 | 19880822 |
| | ES 2067461 | T3 | 19950401 | ES 1988-113617 | 19880822 |
| | KR 9701518 | B1 | 19970211 | KR 1988-10803 | 19880825 |
| PRAI | JP 1987-209214 | A | 19870825 | | |
| | JP 1988-96770 | A | 19880421 | | |
| | JP 1988-206455 | A | 19880822 | | |

L84 ANSWER 18 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1989:592253 CAPLUS
DN 111:192253
TI Temperature-dependent oligomerization of hsp85 in vitro
AU Lanks, Karl W.
CS Health Sci. Cent., SUNY, Brooklyn, NY, 11203, USA
SO Journal of Cellular Physiology (1989), 140(3), 601-7
CODEN: JCLLAX; ISSN: 0021-9541
DT Journal
LA English

L84 ANSWER 19 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1986:439604 CAPLUS
DN 105:39604
TI Phytoalexins, water-**stress** and stomata. III. The effects of some phenolics, fatty acids and some other compounds on stomatal responses
AU Plumbe, Alison M.; Willmer, C. M.
CS Dep. Biol. Sci., Univ. Stirling, Stirling, FK9 4LA, UK
SO New Phytologist (1986), 103(1), 17-22
CODEN: NEPHAV; ISSN: 0028-646X
DT Journal
LA English

L84 ANSWER 20 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1986:183503 CAPLUS
DN 104:183503
TI Phytoalexins, water-**stress** and stomata. II. The effects of phytoalexins on stomatal responses in epidermal strips and on guard cell protoplasts
AU Plumbe, Alison M.; Willmer, C. M.
CS Dep. Biol. Sci., Univ. Stirling, Stirling, FK9 4LA, UK
SO New Phytologist (1986), 102(3), 375-84
CODEN: NEPHAV; ISSN: 0028-646X

DT Journal
LA English

L84 ANSWER 21 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1983:447553 CAPLUS
DN 99:47553
TI Effect of synthetic acyclic polyisoprenoids on the cold-restraint **stress** induced gastric ulcer in rats
AU Murakami, Manabu; Oketani, Kiyoshi; Fujisaki, Hideaki; Wakabayashi, Tsuneo; Inai, Yuichi; Abe, Shinya; Yamatsu, Isao; Ohgo, Toshiharu
CS Tsukuba Res. Lab., Eisai Co., Ltd., Ibaraki, 300-26, Japan
SO Japanese Journal of Pharmacology (1983), 33(3), 549-56
CODEN: JJPAAZ; ISSN: 0021-5198
DT Journal
LA English

L84 ANSWER 22 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1981:12921 CAPLUS
DN 94:12921
TI Effect of abscisic acid on rishitin and lubimin accumulation and resistance to Phytophthora infestans and Cladosporium cucumerinum in potato tuber tissue slices
AU Henfling, J. W. D. M.; Bostock, R.; Kuc, J.
CS Dep. Plant Pathol., Univ. Kentucky, Lexington, KY, 40546, USA
SO Phytopathology (1980), 70(11), 1074-8
CODEN: PHYTAJ; ISSN: 0031-949X
DT Journal
LA English

L84 ANSWER 23 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1979:51352 CAPLUS
DN 90:51352
TI The role of abscisic acid and farnesol in the alleviation of water **stress**
AU Mansfield, T. A.; Wellburn, A. R.; Moreira, T. J. S.
CS Dep. Biol. Sci., Univ. Lancaster, Bailrigg/Lancaster, UK
SO Philosophical Transactions of the Royal Society of London, Series B: Biological Sciences (1978), 284(1002), 471-82
CODEN: PTRBAE; ISSN: 0080-4622
DT Journal; General Review
LA English

L84 ANSWER 24 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1977:597601 CAPLUS
DN 87:197601
TI The role of farnesol as a regulator of stomatal opening in Sorghum
AU Fenton, R.; Davies, W. J.; Mansfield, T. A.
CS Dep. Biol. Sci., Univ. Lancaster, Bailrigg/Lancaster, UK
SO Journal of Experimental Botany (1977), 28(105), 1043-53
CODEN: JEBOA6; ISSN: 0022-0957
DT Journal
LA English

L84 ANSWER 25 OF 26 CAPLUS COPYRIGHT 2003 ACS
AN 1975:121655 CAPLUS
DN 82:121655
TI All-trans-farnesol. Naturally occurring antitranspirant
AU Wellburn, A. R.; Ogunkanmi, A. B.; Fenton, R.; Mansfield, T. A.
CS Dep. Biol. Sci., Univ. Lancaster, Lancaster, UK
SO Planta (1974), 120(3), 255-63
CODEN: PLANAB; ISSN: 0032-0935
DT Journal

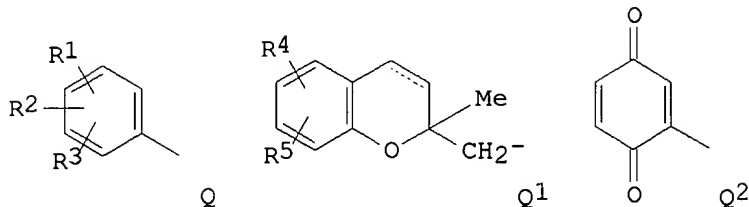
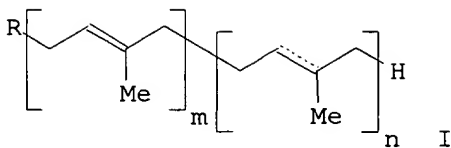
LA English
 L84 ANSWER 26 OF 26 CAPLUS COPYRIGHT 2003 ACS
 AN 1974:532796 CAPLUS
 DN 81:132796
 TI Detection and preliminary identification of endogenous antitranspirants in water-stressed sorghum plants
 AU Ogunkanmi, A. B.; Wellburn, A. R.; Mansfield, T. A.
 CS Dep. Biol. Sci., Univ. Lancaster, Lancaster, UK
 SO Planta (1974), 117(4), 293-302
 CODEN: PLANAB; ISSN: 0032-0935
 DT Journal
 LA English

=> d 184 17 all

L84 ANSWER 17 OF 26 CAPLUS COPYRIGHT 2003 ACS
 AN 1990:36226 CAPLUS
 DN 112:36226
 TI Preparation of antiulcer isoprenoid derivatives and pharmaceutical compositions containing them
 IN Yamada, Kazuhiko; Tahara, Yoshiyuki; Toyoda, Masashi; Irino, Osamu; Misaki, Noriyuki
 PA Nisshin Flour Milling Co., Ltd., Japan
 SO Eur. Pat. Appl., 29 pp.
 CODEN: EPXXDW
 DT Patent
 LA English
 IC ICM A61K031-05
 ICS A61K031-085; A61K031-09; A61K031-22; A61K031-12; A61K031-355;
 C07C050-06; C07D311-72; C07D311-58; C07C039-19
 CC 30-40 (Terpenes and Terpenoids)
 Section cross-reference(s): 1
 FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|---|------|----------|-----------------|----------|
| PI | EP 304842 | A2 | 19890301 | EP 1988-113617 | 19880822 |
| | EP 304842 | A3 | 19910116 | | |
| | EP 304842 | B1 | 19941130 | | |
| | R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE | | | | |
| | JP 02042030 | A2 | 19900213 | JP 1988-206465 | 19880822 |
| | US 4906669 | A | 19900306 | US 1988-234895 | 19880822 |
| | ES 2067461 | T3 | 19950401 | ES 1988-113617 | 19880822 |
| | KR 9701518 | B1 | 19970211 | KR 1988-10803 | 19880825 |
| PRAI | JP 1987-209214 | A | 19870825 | | |
| | JP 1988-96770 | A | 19880421 | | |
| | JP 1988-206455 | A | 19880822 | | |

GI



- AB The title compds. [I; R = Q, Q1, Q2; R1-R3 = H, OH, alkanoyloxy, alkyl, alkoxy, provided that .gtoreq.2 .noteq. H; R4, R5 = H, OH, alkanoyloxy; m = 0, 1; n = 0-9], useful as antiulcer agents, were prep'd. Hydroquinone in dioxane contg. BF3.Et2O was heated 1 h at 40.degree. with 3,7,11,15-tetramethyl-2-hexadecen-1-ol to give, after hydrolysis, 2-(3,7,11,15-tetramethyl-2-hexadecenyl)hydroquinone. At 100 mg/kg orally the latter gave 84.9% inhibition of **stress**-induced ulcers in male rats. A tablet was formulated contg. 2-[3,7(R),11(R),15-tetramethyl-2-hexadecenyl]hydroquinone 50, cryst. cellulose 50, cornstarch 30, lactose 17.8, hydroxypropylcellulose 0.2, and Mg stearate 2 mg.
- ST isoprenoid deriv prep'n antiulcer; phytolhydroquinone deriv prep'n antiulcer; hydroquinone phytol prep'n antiulcer
- IT Ulcer inhibitors
(isoprenoid derivs.)
- IT 50848-64-1P 61977-06-8P 71258-98-5P 77551-14-5P 79577-58-5P
102043-79-8P 123086-31-7P 123086-32-8P
- RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
(prep'n. and reaction of, in prep'n. of ulcer inhibitors)
- IT 119-98-2P 6199-76-4P 10457-66-6P 35175-62-3P 39703-09-8P
71258-97-4P 74166-02-2P 74166-03-3P 76275-68-8P 81557-69-9P
113714-90-2P 119980-00-6P 123086-31-7P 123086-32-8P 123086-33-9P
123086-34-0P 123086-35-1P 123086-36-2P 123086-37-3P 123086-38-4P
123086-39-5P 123086-40-8P 123086-41-9P 123086-42-0P 123086-43-1P
123086-44-2P 123086-45-3P 123086-46-4P 123086-47-5P 123086-48-6P
123086-49-7P 123086-50-0P 123086-51-1P 123086-52-2P 123086-53-3P
123086-54-4P 123086-55-5P 123086-56-6P 123086-57-7P 123086-58-8P
123086-59-9P 123086-60-2P 123086-61-3P 123086-62-4P 123086-63-5P
123086-64-6P 123164-53-4P 123237-20-7P
- RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
(prep'n. of, as ulcer inhibitor)
- IT 87-66-1, 1,2,3-Benzenetriol 91-16-7, 1,2-Dimethoxybenzene 106-24-1, Geraniol 119-98-2 120-80-9, 1,2-Benzenediol, reactions 123-31-9, 1,4-Benzenediol, reactions 128-39-2, 2,6-Di-tert-butylphenol 141-97-9, Ethyl acetoacetate 1113-21-9, Geranylinalool **4602-84-0**, Farnesol 7541-49-3 **13190-97-1**, Solanesol 77551-14-5 79577-58-5 123086-37-3 123086-40-8 123086-45-3 123086-47-5 123164-54-5
- RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, in prep'n. of ulcer inhibitors)

=> s 181 and 119

L85 6 L81 AND L19

=> d 185 1-6

L85 ANSWER 1 OF 6 CAPLUS COPYRIGHT 2003 ACS

AN 2001:923556 CAPLUS

DN 136:58521

TI Cosmetic composition for stressed skin under extreme conditions containing a hydrocarbon, a silicone and plant extracts

IN Mohammadi, Fatemeh; Vargas, Anthony

PA FD Management, Inc., USA

SO PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|------|-----------------|------|
|------------|------|------|-----------------|------|

| | | | | |
|------------------|----|----------|-----------------|----------|
| PI WO 2001095728 | A1 | 20011220 | WO 2001-US19200 | 20010613 |
|------------------|----|----------|-----------------|----------|

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ,
VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

| | | | | |
|---------------|----|----------|----------------|----------|
| US 2002012640 | A1 | 20020131 | US 2001-880245 | 20010613 |
|---------------|----|----------|----------------|----------|

PRAI US 2000-211290P P 20000613

RE.CNT 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 2 OF 6 CAPLUS COPYRIGHT 2003 ACS

AN 2001:137173 CAPLUS

DN 134:178396

TI Synthesis, activity and formulations of pharmaceutical compounds for treatment of oxidative **stress** and/or endothelial dysfunction

IN Del Soldato, Piero

PA Nicox S.A., Fr.

SO PCT Int. Appl., 94 pp.

CODEN: PIXXD2

DT Patent

LA English

FAN.CNT 1

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|------|------|-----------------|------|
|------------|------|------|-----------------|------|

| | | | | |
|------------------|----|----------|----------------|----------|
| PI WO 2001012584 | A2 | 20010222 | WO 2000-EP7225 | 20000727 |
|------------------|----|----------|----------------|----------|

| | | | | |
|---------------|----|----------|--|--|
| WO 2001012584 | A3 | 20020829 | | |
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HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG,
MK, MN, MX, NO, NZ, PL, RO, SG, SI, SK, TR, TT, UA, US, UZ, VN,
YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG

| | | | | |
|---------------|---|----------|---------------|----------|
| BR 2000013264 | A | 20020416 | BR 2000-13264 | 20000727 |
|---------------|---|----------|---------------|----------|

| | | | | |
|------------|----|----------|----------------|----------|
| EP 1252133 | A2 | 20021030 | EP 2000-953102 | 20000727 |
|------------|----|----------|----------------|----------|

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO, MK, CY, AL

| | | | | |
|---------------|----|----------|----------------|----------|
| JP 2003515526 | T2 | 20030507 | JP 2001-516885 | 20000727 |
|---------------|----|----------|----------------|----------|

NO 2002000623 A 20020409 NO 2002-623 20020208
 PRAI IT 1999-MI1817 A 19990812
 WO 2000-EP7225 W 20000727
 OS MARPAT 134:178396

L85 ANSWER 3 OF 6 CAPLUS COPYRIGHT 2003 ACS
 AN 2000:742057 CAPLUS
 DN 133:309791
 TI Synthesis, activity and formulations of pharmaceutical compounds for treatment of oxidative **stress** and/or endothelial dysfunction
 IN Del Soldato, Piero
 PA Nicox S.A., Fr.
 SO PCT Int. Appl., 140 pp.
 CODEN: PIXXD2
 DT Patent
 LA English
 FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|---|------------|----------|-----------------|----------|
| PI | WO 2000061541 | A2 | 20001019 | WO 2000-EP3239 | 20000411 |
| | WO 2000061541 | A3 | 20010927 | | |
| | W: AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, DM, EE, GE, HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG, MK, MN, MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM | | | | |
| | RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG | | | | |
| | IT 1311923 | B1 | 20020320 | IT 1999-MI752 | 19990413 |
| | BR 2000009703 | A | 20020108 | BR 2000-9703 | 20000411 |
| | EP 1169298 | A2 | 20020109 | EP 2000-926870 | 20000411 |
| | R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO | | | | |
| | JP 2002541236 | T2 | 20021203 | JP 2000-610818 | 20000411 |
| | NO 2001004928 | A | 20011213 | NO 2001-4928 | 20011010 |
| PRAI | IT 1999-MI752 | A | 19990413 | | |
| | WO 2000-EP3239 | W | 20000411 | | |
| OS | MARPAT | 133:309791 | | | |

L85 ANSWER 4 OF 6 CAPLUS COPYRIGHT 2003 ACS
 AN 2000:742053 CAPLUS
 DN 133:310142
 TI Synthesis, activity and formulations of pharmaceutical compounds for treatment of oxidative **stress** and/or endothelial dysfunction
 IN Del Soldato, Piero
 PA Nicox S.A., Fr.
 SO PCT Int. Appl., 159 pp.
 CODEN: PIXXD2
 DT Patent
 LA English
 FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----|---|------|----------|-----------------|----------|
| PI | WO 2000061537 | A2 | 20001019 | WO 2000-EP3234 | 20000411 |
| | WO 2000061537 | A3 | 20010927 | | |
| | W: AL, AU, BA, BB, BG, BR, CA, CN, CU, CZ, DM, EE, GE, HR, HU, ID, IL, IN, IS, JP, KP, KR, LC, LK, LR, LT, LV, MA, MG, MK, MN, MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM | | | | |
| | RW: GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG | | | | |

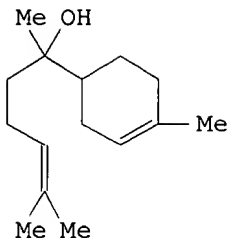
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|--|-------------------|----------|----------------|----------|
| IT 1311924 | B1 | 20020320 | IT 1999-MI753 | 19990413 |
| BR 2000009702 | A | 20020108 | BR 2000-9702 | 20000411 |
| EP 1169294 | A2 | 20020109 | EP 2000-925203 | 20000411 |
| R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, SI, LT, LV, FI, RO | | | | |
| JP 2002541233 | T2 | 20021203 | JP 2000-610814 | 20000411 |
| NO 2001004927 | A | 20011213 | NO 2001-4927 | 20011010 |
| PRAI | IT 1999-MI753 | A | 19990413 | |
| | WO 2000-EP3234 | W | 20000411 | |
| OS | MARPAT 133:310142 | | | |

L85 ANSWER 5 OF 6 CAPLUS COPYRIGHT 2003 ACS
AN 1999:168599 CAPLUS
DN 130:335402
TI Chemical response of parsley and Mentha herbs to certain **stress** agents
AU Hashema, Fatma Abd El-Megeed; Sahab, Ahmed Farahat
CS Pharmaceutical Science Department, National Research Centre, Cairo, Egypt
SO Food Chemistry (1999), 65(1), 29-33
CODEN: FOCHDJ; ISSN: 0308-8146
PB Elsevier Science Ltd.
DT Journal
LA English
RE.CNT 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD
ALL CITATIONS AVAILABLE IN THE RE FORMAT

L85 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2003 ACS
AN 1979:400383 CAPLUS
DN 91:383
TI Pharmacological experiments with components of chamomile. III.
Experimental animal studies of the ulcer-protective effect of chamomile
AU Szelényi, I.; Isaac, O.; Thiemer, K.
CS Chemiewerk Homburg, Degussa, Frankfurt/Main, Fed. Rep. Ger.
SO Planta Medica (1979), 35(2), 218-27
CODEN: PLMEAA; ISSN: 0032-0943
DT Journal
LA German

=> d 185 6 all

L85 ANSWER 6 OF 6 CAPLUS COPYRIGHT 2003 ACS
AN 1979:400383 CAPLUS
DN 91:383
TI Pharmacological experiments with components of chamomile. III.
Experimental animal studies of the ulcer-protective effect of chamomile
AU Szelényi, I.; Isaac, O.; Thiemer, K.
CS Chemiewerk Homburg, Degussa, Frankfurt/Main, Fed. Rep. Ger.
SO Planta Medica (1979), 35(2), 218-27
CODEN: PLMEAA; ISSN: 0032-0943
DT Journal
LA German
CC 1-5 (Pharmacodynamics)
GI



AB (-)-.alpha.-Bisabolol (I) [23089-26-1], a component of
 chamomile, inhibited the ulcer formation induced by indomethacin, alc., or
stress in rats, and increased the rate of healing of ulcers caused
 by HOAc or heat cauterization of the stomach. The chamomile ext.
 Kamillosan also inhibited the occurrence of alc.-induced ulceration.
 ST bisabolol ulcer inhibition; chamomile component ulcer inhibition
 IT Ulcer
 (bisabolol and chamomile ext. inhibition of)
 IT Chamomile
 (ext. of, ulcer inhibition by)
 IT 23089-26-1
 RL: BIOL (Biological study)
 .ulcer inhibition by)

=> s 121 and 181
 L86 2 L21 AND L81

=> s 186 1-2
 MISSING OPERATOR L86 1-2
 The search profile that was entered contains terms or
 nested terms that are not separated by a logical operator.

=> d 186 1-2

L86 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS
 AN 2002:392183 CAPLUS
 DN 136:385060
 TI Biooxidation of volatile organics by Candida sp.
 IN Eirich, L. Dudley; Anderson, Kevin W.; Gates, Jeffrey A.; Wilson, C. Ron;
 Biermann, Manfred; Vice, Gilbert H.
 PA USA
 SO U.S. Pat. Appl. Publ., 17 pp.
 CODEN: USXXCO
 DT Patent
 LA English
 FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|-------------------|------|----------|-----------------|----------|
| PI | US 2002061566 | A1 | 20020523 | US 2001-812308 | 20010320 |
| PRAI | US 2000-190626P | P | 20000320 | | |
| OS | MARPAT 136:385060 | | | | |

L86 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS
 AN 1990:191786 CAPLUS
 DN 112:191786
 TI Measures of anxiety, retention and **stress** in the rat following
 treatment with the diterpene sclareol glycol
 AU Georgieva, Zh.
 CS Inst. Pharmacol. Pharm., Sofia, 1431, Bulg.
 SO Methods and Findings in Experimental and Clinical Pharmacology (1990),

12(1), 5-10
CODEN: MFEPPDX; ISSN: 0379-0355
DT Journal
LA English

=> d 186 2 all

L86 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS
AN 1990:191786 CAPLUS
DN 112:191786
TI Measures of anxiety, retention and **stress** in the rat following treatment with the diterpene sclareol glycol
AU Georgieva, Zh.
CS Inst. Pharmacol. Pharm., Sofia, 1431, Bulg.
SO Methods and Findings in Experimental and Clinical Pharmacology (1990), 12(1), 5-10
CODEN: MFEPPDX; ISSN: 0379-0355
DT Journal
LA English
CC 1-11 (Pharmacology)
Section cross-reference(s): 14
AB In a punished drinking test in rats sclareol glycol (SG) decreased the no. of punished responses (proconflict response) while diazepam had the opposite effect; SG antagonized the anticonflict response of diazepam. Post-training administration of SG in rats enhanced retention in active avoidance task evaluated 24 h later. SG produced an increase in plasma ACTH and corticosterone levels in unstressed rats. The **stress**-induced increase in ACTH and corticosterone secretion was potentiated by SG. These data suggest that SG behaves as an anxiogenic, memory-facilitator and perhaps adaptogenic agent. The effects of SG may be mediated by different mechanisms of action (stimulation of adenylate cyclase or interaction with GABA-ergic and dopaminergic transmitter mechanisms).
ST sclareol glycol anxiety learning **stress**; diterpene anxiety learning **stress**
IT **Stress**, biological
(ACTH and corticosterone secretion induction by, sclareol glycol enhancement of)
IT Anxiety
(from sclareol glycol)
IT Learning
(sclareol glycol enhancement of)
IT 38419-75-9, Sclareol glycol
RL: BIOL (Biological study)
(anxiety from and learning stimulation by and **stress**-induced increase in ACTH and corticosterone secretion response to)
IT 50-22-6, Corticosterone 9002-60-2, ACTH, biological studies
RL: BIOL (Biological study)
(secretion of, sclareol glycol increase of **stress**-induced)

=> s 174 and 181
L87 1 L74 AND L81

=> s 187
L88 1 L74 AND L81

=> d 187

L87 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
AN 1990:36226 CAPLUS

DN 112:36226
 TI Preparation of antiulcer isoprenoid derivatives and pharmaceutical compositions containing them
 IN Yamada, Kazuhiko; Tahara, Yoshiyuki; Toyoda, Masashi; Irino, Osamu; Misaki, Noriyuki
 PA Nisshin Flour Milling Co., Ltd., Japan
 SO Eur. Pat. Appl., 29 pp.
 CODEN: EPXXDW

DT Patent

LA English

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | |
|----|---|----------------|----------|-----------------|----------|--|
| PI | EP 304842 | A2 | 19890301 | EP 1988-113617 | 19880822 | |
| | EP 304842 | A3 | 19910116 | | | |
| | EP 304842 | B1 | 19941130 | | | |
| | R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE | | | | | |
| | JP 02042030 | A2 | 19900213 | JP 1988-206465 | 19880822 | |
| | US 4906669 | A | 19900306 | US 1988-234895 | 19880822 | |
| | ES 2067461 | T3 | 19950401 | ES 1988-113617 | 19880822 | |
| | KR 9701518 | B1 | 19970211 | KR 1988-10803 | 19880825 | |
| | PRAI | JP 1987-209214 | A | 19870825 | | |
| | | JP 1988-96770 | A | 19880421 | | |
| | JP 1988-206455 | A | 19880822 | | | |

=> s 187 1 all

MISSING OPERATOR L87 1 ALL

The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

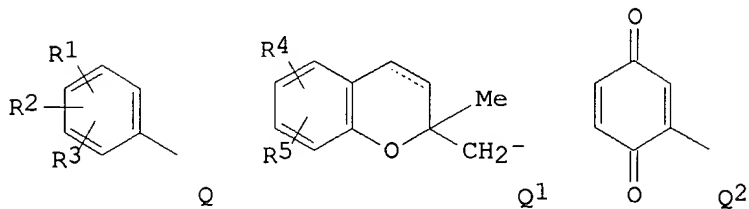
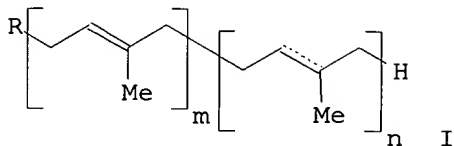
=> d 187 1 all

L87 ANSWER 1 OF 1 CAPLUS COPYRIGHT 2003 ACS
 AN 1990:36226 CAPLUS
 DN 112:36226
 TI Preparation of antiulcer isoprenoid derivatives and pharmaceutical compositions containing them
 IN Yamada, Kazuhiko; Tahara, Yoshiyuki; Toyoda, Masashi; Irino, Osamu; Misaki, Noriyuki
 PA Nisshin Flour Milling Co., Ltd., Japan
 SO Eur. Pat. Appl., 29 pp.
 CODEN: EPXXDW
 DT Patent
 LA English
 IC ICM A61K031-05
 ICS A61K031-085; A61K031-09; A61K031-22; A61K031-12; A61K031-355; C07C050-06; C07D311-72; C07D311-58; C07C039-19
 CC 30-40 (Terpenes and Terpenoids)
 Section cross-reference(s): 1
 FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | |
|----|---|----------------|----------|-----------------|----------|--|
| PI | EP 304842 | A2 | 19890301 | EP 1988-113617 | 19880822 | |
| | EP 304842 | A3 | 19910116 | | | |
| | EP 304842 | B1 | 19941130 | | | |
| | R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE | | | | | |
| | JP 02042030 | A2 | 19900213 | JP 1988-206465 | 19880822 | |
| | US 4906669 | A | 19900306 | US 1988-234895 | 19880822 | |
| | ES 2067461 | T3 | 19950401 | ES 1988-113617 | 19880822 | |
| | KR 9701518 | B1 | 19970211 | KR 1988-10803 | 19880825 | |
| | PRAI | JP 1987-209214 | A | 19870825 | | |
| | | | | | | |

JP 1988-96770 A 19880421
JP 1988-206455 A 19880822

GI



AB The title compds. [I; R = Q, Q1, Q2; R1-R3 = H, OH, alkanoyloxy, alkyl, alkoxy, provided that $\text{gtoreq} 2 \text{ .noteq. } \text{H}$; R4, R5 = H, OH, alkanoyloxy; m = 0, 1; n = 0-9], useful as antiulcer agents, were prep'd. Hydroquinone in dioxane contg. BF3.Et2O was heated 1 h at 40.degree. with 3,7,11,15-tetramethyl-2-hexadecen-1-ol to give, after hydrolysis, 2-(3,7,11,15-tetramethyl-2-hexadecenyl)hydroquinone. At 100 mg/kg orally the latter gave 84.9% inhibition of stress-induced ulcers in male rats. A tablet was formulated contg. 2-[3,7(R),11(R),15-tetramethyl-2-hexadecenyl]hydroquinone 50, cryst. cellulose 50, cornstarch 30, lactose 17.8, hydroxypropylcellulose 0.2, and Mg stearate 2 mg.

ST isoprenoid deriv prep'n antiulcer; phytolhydroquinone deriv prep'n antiulcer; hydroquinone phytol prep'n antiulcer

IT Ulcer inhibitors

(isoprenoid derivs.)

IT 50848-64-1P 61977-06-8P 71258-98-5P 77551-14-5P 79577-58-5P
102043-79-8P 123086-31-7P 123086-32-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prep'n. and reaction of, in prep'n. of ulcer inhibitors)

IT 119-98-2P 6199-76-4P 10457-66-6P 35175-62-3P 39703-09-8P
71258-97-4P 74166-02-2P 74166-03-3P 76275-68-8P 81557-69-9P
113714-90-2P 119980-00-6P 123086-31-7P 123086-32-8P 123086-33-9P
123086-34-0P 123086-35-1P 123086-36-2P 123086-37-3P 123086-38-4P
123086-39-5P 123086-40-8P 123086-41-9P 123086-42-0P 123086-43-1P
123086-44-2P 123086-45-3P 123086-46-4P 123086-47-5P 123086-48-6P
123086-49-7P 123086-50-0P 123086-51-1P 123086-52-2P 123086-53-3P
123086-54-4P 123086-55-5P 123086-56-6P 123086-57-7P 123086-58-8P
123086-59-9P 123086-60-2P 123086-61-3P 123086-62-4P 123086-63-5P
123086-64-6P 123164-53-4P 123237-20-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(prep'n. of, as ulcer inhibitor)

IT 87-66-1, 1,2,3-Benzenetriol 91-16-7, 1,2-Dimethoxybenzene 106-24-1,
Geraniol 119-98-2 120-80-9, 1,2-Benzenediol, reactions 123-31-9,
1,4-Benzenediol, reactions 128-39-2, 2,6-Di-tert-butylphenol 141-97-9,
Ethyl acetoacetate 1113-21-9, Geranylinalool 4602-84-0,

Farnesol 7541-49-3 13190-97-1, Solanesol 77551-14-5 79577-58-5
123086-37-3 123086-40-8 123086-45-3 123086-47-5 123164-54-5
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, in prepn. of ulcer inhibitors)

=> d his

(FILE 'HOME' ENTERED AT 15:49:40 ON 15 JUL 2003)

FILE 'REGISTRY' ENTERED AT 15:49:46 ON 15 JUL 2003

L1 43 S CEDROL
L2 33 S PATCHOULI
L3 36 S SANTALOL
L4 31 S BISABOLOL
E BISABOLOL
L5 2 S VETIVEROL
L6 30 S SCLAREOL
L7 0 S GLOBUOL
L8 9 S GLOBULOL
L9 11 S GUAIOL

FILE 'CAPLUS' ENTERED AT 15:55:44 ON 15 JUL 2003

E SLEEP
L10 15616 S E3
E SEDATIVE
L11 12852 S E3-E9
E RELAXATION
L12 217668 S E3 OR E8
E NARCOTIC
L13 10398 S E3 OR E12
E HYPNOTIC
L14 9427 S E3 OR E10
E SOMULENCE
E SOMUL
E SOMNIA
E INSOMNIA
L15 1431 S E3-E8
L16 717 S L1
L17 214 S L2
L18 350 S L3
L19 1195 S L4
L20 80 S L5
L21 487 S L6
L22 621 S L8
L23 12852 S L11
L24 12852 S L11
L25 422 S L9
L26 1 S L6 AND L10

FILE 'REGISTRY' ENTERED AT 16:04:01 ON 15 JUL 2003

L27 1 S 38419-75-9/RN
SET NOTICE 1 DISPLAY
SET NOTICE LOGIN DISPLAY

FILE 'CAPLUS' ENTERED AT 16:04:27 ON 15 JUL 2003

L28 2 S L10 AND L16
L29 1 S L16 AND L11
L30 0 S L16 AND L12
L31 0 S L16 AND L13
L32 1 S L16 AND L14
L33 0 S L15 AND L16

L34 0 S L17 AND L10
L35 0 S L17 AND L11
L36 0 S L17 AND L13
L37 0 S L17 AND L15
L38 0 S L18 AND L10
L39 0 S L18 AND L13
L40 0 S L19 AND L10
L41 0 S LL18 AND L12
L42 0 S L18 AND L12
L43 1 S L19 AND L12
L44 1 S 43 1 ALL
L45 0 S L19 AND L15
L46 0 S L20 AND L10
L47 0 S L20 AND L12
L48 1 S L21 AND L10
L49 0 S L22 AND L12
L50 0 S L25 AND L10
E NERVIOS
E NERVOUS
L51 164186 S E3-E7
L52 1 S L51 AND L16

FILE 'REGISTRY' ENTERED AT 16:16:22 ON 15 JUL 2003

E FARNESOL
L53 75 S E3
E EUGENOL
L54 165 S E3
L55 4 S GERANYL LINALOOL
E CEDRENOL
L56 9 S E3
E ISOPYTOL
E ISOPHYTOL
L57 6 S E3
E NEROLIDOL
L58 41 S E3

FILE 'CAPLUS' ENTERED AT 16:21:24 ON 15 JUL 2003

1 S L56 AND L10
L60 0 S L56 AND L12
L61 0 S L56 AND L15
L62 90 S L56
L63 1 S L62 AND L10
L64 0 S L62 AND L12
L65 3275 S L53
L66 1 S L65 AND L10
L67 9 S L65 AND L12
L68 0 S L65 AND L13
L69 0 S L65 AND L15
L70 8961 S L54
L71 8 S L70 AND L10
L72 33 S L70 AND L12
L73 33 S L72 NOT L71
L74 123 S L55
L75 1 S L74 AND L10
L76 380 S L57
L77 1 S L76 AND L10
L78 0 S L76 AND L15
L79 2587 S L58
L80 2 S L79 AND L10
E STRESS
L81 393704 S E3
L82 5 S L81 AND L16

L83 0 S L62 AND L81
L84 26 S L65 AND L81
L85 6 S L81 AND L19
L86 2 S L21 AND L81
L87 1 S L74 AND L81
L88 1 S L87

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---Logging off of STN---

=>

Executing the logoff script...

=> LOG Y

| COST IN U.S. DOLLARS | SINCE FILE ENTRY | TOTAL SESSION |
|----------------------|------------------|---------------|
| FULL ESTIMATED COST | 131.09 | 336.77 |

| DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) | SINCE FILE ENTRY | TOTAL SESSION |
|--|------------------|---------------|
| CA SUBSCRIBER PRICE | -7.81 | -10.41 |

STN INTERNATIONAL LOGOFF AT 16:46:11 ON 15 JUL 2003

AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian
CC 1-11 (Pharmacology)
AB In expts. on male mice and rats some behavioral effects of the diterpene sclareol glucol (SG) were studied. SG in higher doses (50, 100 mg/kg) enhanced the sensitivity to touch and pain, shortened the duration of hexobarbital **sleep**, stimulated exploratory behavior in rats in open field, exerted no anti-convulsive effect and increased the mortality from pentylenetetrazole seizures. In higher doses (300, 500, and 1000 mg/kg) SG induced seizures.
ST sclareol glucol behavior nervous system
IT Behavior
Nervous system
(clareol glucol effect on)
IT **38419-75-9**
RL: PRP (Properties)
(behavioral and nervous systems effects of)

AN 1992:171087 CAPLUS
DN 116:171087
TI Effects of olfactory stimulation with jasmin and its component chemicals on the duration of pentobarbital-induced **sleep** in mice
AU Tsuchiya, T.; Tanida, M.; Uenoyama, S.; Nakayama, Y.
CS Shiseido Res. Cent., Yokohama, 223, Japan
SO Life Sciences (1992), 50(15), 1097-102
CODEN: LIFSAK; ISSN: 0024-3205
DT Journal
LA English
CC 13-6 (Mammalian Biochemistry)
Section cross-reference(s): 62
AB The effect of olfactory stimulation with jasmin and its component chems. on pentobarbital **sleep** time was investigated using mice in order to det. which component of jasmin influences pentobarbital **sleep** time via olfactory stimulation. **Sleep** time was defined as the time elapsed between i.p. pentobarbital administration and the first time that the animal was able to spontaneously right itself. **Sleep** time was significantly decreased by olfactory stimulation with jasmin, and also by one of the fractions obtained by fractional distn. at 150 .degree.C and 0.1 mmHg. The fraction which influenced the **sleep** time was found to consist of benzyl benzoate, isophytol, geranyl linalool, phytol and phytol acetate, which were identified using gas chromatog. with mass and IR spectrometry. In expts. using authentic samples of these components, phytol significantly shortened the pentobarbital **sleep** time, while the others had no effect. Phytol is the component of jasmin which reduces the duration of pentobarbital-induced **sleep**.
ST **sleep** pentobarbital jasmin phytol drug interaction; olfactory system **sleep** pentobarbital jasmin phytol
IT **sleep**
 (jasmin inhibition of pentobarbital-induced, olfactory stimulation in)
IT Essential oils
 RL: BIOL (Biological study)
 (jasmine, Jasminum grandiflorum abs., pentobarbital-induced **sleep** inhibition by, olfactory stimulation in)
IT Nervous system
 (olfactory system, jasmin stimulation of, pentobarbital-induced **sleep** inhibition by)
IT 76-74-4, Pentobarbital
 RL: BIOL (Biological study)
 (jasmin inhibition of **sleep** stimulation by, olfactory stimulation in)
IT 120-51-4, Benzyl benzoate 140-11-4, Benzyl acetate 150-86-7, Phytol 505-32-8, Isophytol 1113-21-9, Geranyl linalool 10236-16-5, Phytol acetate
 RL: BIOL (Biological study)
 (pentobarbital **sleep** time response to, as jasmin component, olfactory stimulation in relation to)

=>

AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian
CC 1-11 (Pharmacology)
AB In expts. on male mice and rats some behavioral effects of the diterpene sclareol glucol (SG) were studied. SG in higher doses (50, 100 mg/kg) enhanced the sensitivity to touch and pain, shortened the duration of hexobarbital **sleep**, stimulated exploratory behavior in rats in open field, exerted no anti-convulsive effect and increased the mortality from pentylenetetrazole seizures. In higher doses (300, 500, and 1000 mg/kg) SG induced seizures.
ST sclareol glucol behavior nervous system
IT Behavior
Nervous system
(sclareol glucol effect on)
IT **38419-75-9**
RL: PRP (Properties)
(behavioral and nervous systems effects of)

AN 1993:616760 CAPLUS
DN 119:216760
TI Calcium antagonistic properties of the sesquiterpene T-cadinol and related substances: structure-activity studies
AU Zygmunt, P. M.; Larsson, B.; Sterner, O.; Vinge, E.; Hoegestaett, E. D.
CS Dep. Clin. Pharmacol., Univ. Hosp. Lund, Lund, S-22185, Swed.
SO Pharmacology & Toxicology (Oxford, United Kingdom) (1993), 73(1), 3-9
CODEN: PHTOEH; ISSN: 0901-9928
DT Journal
LA English
CC 1-3 (Pharmacology)
AB The calcium antagonistic properties of (+)-T-cadinol, some of its stereoisomers and related terpenes were investigated in both functional and radioligand binding studies, and the effects were compared with those of the dihydropyridine calcium antagonist (.+-.)-nimodipine. In the isolated rat aorta, the terpenes relaxed contractions induced by 60 mM K⁺ more potently than those induced by phenylephrine. (+)-T-cadinol and its stereoisomers were the most potent among the terpenes to relax K⁺-induced contractions, whereas they were approx. 10,000 times less potent than (.+-.)-nimodipine in this regard. Binding of the dihydropyridine radioligand [3H]-(+)-PN200-110 was studied on rat cerebral cortical membranes. Displacement and satn. studies indicated that (+)-T-cadinol caused a competitive inhibition of binding. The log Ki values for (+)-T-cadinol and (.+-.)-nimodipine from displacement studies (-4.7 and -9.2) corresponded with the log RC50 values for relaxation of K⁺-contracted rat aortas (-5.0 and -9.0). For the terpenes, there was a significant correlation ($P < 0.001$, $r_s = 0.89$) between displacement of dihydropyridine binding and the ability to relax K⁺-induced contractions. The structures of three terpenes were chem. modified by blocking hydroxyl groups. The potency of these derivs., as well as the naturally occurring deriv. 2-oxo-T-cadinol, to relax K⁺-induced contractions was not correlated to the lipophilicity of the compds. Instead, other qualities appear to be of importance for the functional effects. The authors' results suggest that (+)-T-cadinol and related terpenes may represent a new chem. class of calcium antagonists, which interact with dihydropyridine binding sites on the voltage-operated calcium channels.
ST calcium antagonist terpene T-cadinol structure
IT Terpenes and Terpenoids, biological studies
RL: BIOL (Biological study)
 (calcium antagonism by, structure in relation to)
IT Lipophilicity
 (of sesquiterpene T-cadinol and related substances, calcium antagonism in relation to)
IT Ion channel blockers
 (calcium, sesquiterpene T-cadinol and related substances as, structure in relation to)
IT Molecular structure-biological activity relationship
 (calcium channel-blocking, of sesquiterpene T-cadinol and related substances)
IT Receptors
RL: BIOL (Biological study)
 (dihydropyridine, sesquiterpene T-cadinol and related substances binding to, calcium antagonism by, structure in relation to)
IT 481-34-5, (-)-alpha.-Cadinol 2216-51-5, (-)-Menthol 5937-11-1, (+)-T-Cadinol 19435-97-3 19912-62-0, (-)-T-Muurolol 23089-26-1, (-)-alpha.-Bisabolol 53402-16-7 74638-12-3, (-)-Furosardonin A 129058-89-5, (-)-Tremediol 150718-45-9 150718-46-0 150718-47-1
RL: BIOL (Biological study)
 (calcium antagonism by, structure in relation to)

=>

AN 1987:611773 CAPLUS
DN 107:211773
TI Behavioral effects of the diterpene sclareol glucol
AU Georgieva, Zh.; Uzunov, P.; Ognyanov, I.; Bantutova, M.
CS Bulg.
SO Problemi na Farmakologiyata (1986), 1, 24-32
CODEN: PRFAE9
DT Journal
LA Russian
CC 1-11 (Pharmacology)
AB In expts. on male mice and rats some behavioral effects of the diterpene sclareol glucol (SG) were studied. SG in higher doses (50, 100 mg/kg) enhanced the sensitivity to touch and pain, shortened the duration of hexobarbital **sleep**, stimulated exploratory behavior in rats in open field, exerted no anti-convulsive effect and increased the mortality from pentylenetetrazole seizures. In higher doses (300, 500, and 1000 mg/kg) SG induced seizures.
ST sclareol glucol behavior nervous system
IT Behavior
Nervous system
(sclareol glucol effect on)
IT 38419-75-9
RL: PRP (Properties)
(behavioral and nervous systems effects of)

AN 1992:171087 CAPLUS
DN 116:171087
TI Effects of olfactory stimulation with jasmin and its component chemicals on the duration of pentobarbital-induced **sleep** in mice
AU Tsuchiya, T.; Tanida, M.; Uenoyama, S.; Nakayama, Y.
CS Shiseido Res. Cent., Yokohama, 223, Japan
SO Life Sciences (1992), 50(15), 1097-102
CODEN: LIFSAK; ISSN: 0024-3205
DT Journal
LA English
CC 13-6 (Mammalian Biochemistry)
Section cross-reference(s): 62
AB The effect of olfactory stimulation with jasmin and its component chems. on pentobarbital **sleep** time was investigated using mice in order to det. which component of jasmin influences pentobarbital **sleep** time via olfactory stimulation. **Sleep** time was defined as the time elapsed between i.p. pentobarbital administration and the first time that the animal was able to spontaneously right itself. **Sleep** time was significantly decreased by olfactory stimulation with jasmin, and also by one of the fractions obtained by fractional distn. at 150 .degree.C and 0.1 mmHg. The fraction which influenced the **sleep** time was found to consist of benzyl benzoate, isophytol, geranyl linalool, phytol and phytol acetate, which were identified using gas chromatog. with mass and IR spectrometry. In expts. using authentic samples of these components, phytol significantly shortened the pentobarbital **sleep** time, while the others had no effect. Phytol is the component of jasmin which reduces the duration of pentobarbital-induced **sleep**.
ST **sleep** pentobarbital jasmin phytol drug interaction; olfactory system **sleep** pentobarbital jasmin phytol
IT **Sleep**
 (jasmin inhibition of pentobarbital-induced, olfactory stimulation in)
IT Essential oils
 RL: BIOL (Biological study)
 (jasmine, Jasminum grandiflorum abs., pentobarbital-induced **sleep** inhibition by, olfactory stimulation in)
IT Nervous system
 (olfactory system, jasmin stimulation of, pentobarbital-induced **sleep** inhibition by)
IT 76-74-4, Pentobarbital
 RL: BIOL (Biological study)
 (jasmin inhibition of **sleep** stimulation by, olfactory stimulation in)
IT 120-51-4, Benzyl benzoate 140-11-4, Benzyl acetate 150-86-7, Phytol 505-32-8, Isophytol(1113-21-9, Geranyl linalool) 10236-16-5, Phytol acetate
 RL: BIOL (Biological study)
 (pentobarbital **sleep** time response to, as jasmin component, olfactory stimulation in relation to)

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AN 1996:287269 CAPLUS
DN 125:1102
TI Synthesis and pharmacological activity of a eugenol derivative
AU Costa, J. A.; Oliveira, R. A. G.; Barbosa, J. M.; Souza Brito, A. R. M.
CS Lab. Tecnologia Farmaceutica, UFPb, Joao Pessoa, Brazil
SO Revista Brasileira de Farmacia (1994), 75(2), 40-5
CODEN: RBFAAH; ISSN: 0370-372X
PB Associacao Brasileira de Farmaceuticos
DT Journal
LA Portuguese
CC 1-11 (Pharmacology)
Section cross-reference(s): 26
AB The aim of this work was the synthesis of a natural pharmacol. active substance. The target compd. could be prepd. by an oxidative coupling reaction involving a starting material also found in nature. Eugenol, an allyl phenol widely used as a dental local anesthetic, was obtained by a soxhlet extn. of cloves oil from Caryophyllum aromaticus. Eugenol, prepd. by purifn. of the crude oil, was dimerized using potassium ferricyanide, giving dehydrodieugenol (DDE), a substance previously isolated from plants. The two phenolic groups were methylated with di-Me sulfate giving di-O-methyldehydrodieugenol (DMDE). Pharmacol. evaluation of DMDE in mice showed that it has a CNS-depressant effect, characterized by general sluggishness of the animal. It potentiated the sleep induced by sodium pentobarbital (which confirms its depressant activity) and also presented an analgesic effect after chem., mech. and thermal nociceptives stimulus. Furthermore, 50% of the exptl. animals were protected against pentylenetetrazol-induced convulsion and survived. These data confirmed the central depressant activity of DMDE.
ST eugenol deriv prepn central nervous depressant; dimethyldehydrodieugenol prepn central nervous depressant
IT Analgesics
Anticonvulsants and Antiepileptics
Nervous system depressants
 Sleep
 (dimethyldehydrodieugenol prepn. and pharmacol. activity)
IT 13417-56-6P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)
 (dimethyldehydrodieugenol prepn. and pharmacol. activity)
IT 97-53-0, Eugenol
RL: RCT (Reactant); RACT (Reactant or reagent)
 (dimethyldehydrodieugenol prepn. and pharmacol. activity)
IT 4433-08-3P, Dehydrodieugenol
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 (dimethyldehydrodieugenol prepn. and pharmacol. activity)

AN 1989:225383 CAPLUS
DN 110:225383
TI Methyl eugenol: laboratory evaluation in animals
AU Barbosa, P. P. P.; Teixeira, J. R. M.; Melo, M. F. B.; Gusmao, Q. M. W. B.
CS Esc. Cienc. Med., Univ. Fed. Alagoas, Alagoas, Brazil
SO Revista Brasileira de Anestesiologia (1988), 38(6), 393-7
CODEN: RBANAV; ISSN: 0034-7094
DT Journal
LA Portuguese
CC 1-11 (Pharmacology)
AB Me Eugenol, an essential oil fraction obtained from *Caryophyllum aromaticus*, caused central depressing effects with significant hypnotic and myorelaxing action in doses of 40 mg/kg, i.p., for rats and 5 mg/kg, i.v., for rabbits and dogs, rapid induction and satisfactory duration of **sleep** (118.4 s and 47.3 min resp.) in rats, and **sleep** time between 9-12 min in dogs. Anesthetic evolution in dogs was satisfactory, followed by rapid recovery and movement. Me eugenol (20 .mu.g/mL) reduced the atrial muscular cardiac force of contraction (50%) in 20 min. The addn. of Me eugenol to isolated rat diaphragm-nerve prepns. produced muscular contraction blockade under direct and indirect stimulation.
ST methyl eugenol hypnotic muscle relaxant
IT Anesthetics
Hypnotics and Sedatives
Muscle relaxants
(Me eugenol)
IT 93-15-2, Methyl eugenol
RL: BIOL (Biological study)
(hypnotic and muscle-relaxant activities of)

AN 1982:504098 CAPLUS
DN 97:104098
TI The pharmacological effects of a ligroin extract of nutmeg (*Myristica fragrans*)
AU Sherry, C. J.; Ray, L. E.; Herron, R. E.
CS Dep. Biol., Texas A and M Univ., College Station, TX, 77843, USA
SO Journal of Ethnopharmacology (1982), 6(1), 61-6
CODEN: JOETD7; ISSN: 0378-8741
DT Journal
LA English
CC 1-11 (Pharmacology)
Section cross-reference(s): 11, 63
AB A ligroin ext. of nutmeg (*Myristica fragrans*) increased the duration of light and deep **sleep** in the young chicken. The presence of trimyristin [555-45-3] tended to increase the effect of the ext. The ext. did not contain detectable amts. of myristicin [607-91-0], elemicin [487-11-6], safrole [94-59-7], or eugenol [97-53-0], which either individually or collectively have been suggested to be the active agents of nutmeg.
ST nutmeg ext pharmacol; psychotropic nutmeg ext
IT Myristica
 (ext. of, compn. and pharmacol. of)
IT Psychotropics
 (nutmeg ext.)
IT 94-59-7 97-53-0 487-11-6 607-91-0
RL: BIOL (Biological study)
 (nutmeg psychotropic activity in relation to)
IT 555-45-3
RL: BIOL (Biological study)
 (nutmeg psychotropic activity potentiation by)

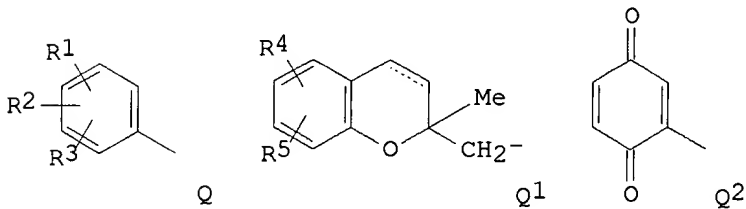
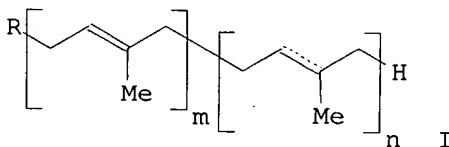
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DN 112:36226
 TI Preparation of antiulcer isoprenoid derivatives and pharmaceutical compositions containing them
 IN Yamada, Kazuhiko; Tahara, Yoshiyuki; Toyoda, Masashi; Irino, Osamu; Misaki, Noriyuki
 PA Nisshin Flour Milling Co., Ltd., Japan
 SO Eur. Pat. Appl., 29 pp.
 CODEN: EPXXDW
 DT Patent
 LA English
 IC ICM A61K031-05
 ICS A61K031-085; A61K031-09; A61K031-22; A61K031-12; A61K031-355;
 C07C050-06; C07D311-72; C07D311-58; C07C039-19
 CC 30-40 (Terpenes and Terpenoids)
 Section cross-reference(s): 1

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------|---|----------|----------------|-----------------|----------|
| PI | EP 304842 | A2 | 19890301 | EP 1988-113617 | 19880822 |
| | EP 304842 | A3 | 19910116 | | |
| | EP 304842 | B1 | 19941130 | | |
| | R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE | | | | |
| | JP 02042030 | A2 | 19900213 | JP 1988-206465 | 19880822 |
| | US 4906669 | A | 19900306 | US 1988-234895 | 19880822 |
| ES 2067461 | T3 | 19950401 | ES 1988-113617 | 19880822 | |
| KR 9701518 | B1 | 19970211 | KR 1988-10803 | 19880825 | |
| PRAI | JP 1987-209214 | A | 19870825 | | |
| | JP 1988-96770 | A | 19880421 | | |
| | JP 1988-206455 | A | 19880822 | | |

GI



AB The title compds. [I; R = Q, Q1, Q2; R1-R3 = H, OH, alkanoyloxy, alkyl, alkoxy, provided that $m \geq 2$ and $n = 0, 1; m = 0-9$], useful as antiulcer agents, were prep'd. Hydroquinone in dioxane contg. BF₃.Et₂O was heated 1 h at 40.degree. with 3,7,11,15-tetramethyl-2-hexadecen-1-ol to give, after hydrolysis, 2-(3,7,11,15-tetramethyl-2-hexadecenyl)hydroquinone. At 100 mg/kg orally the latter gave 84.9% inhibition of stress-induced ulcers in male rats. A tablet was formulated contg. 2-[3,7(R),11(R),15-tetramethyl-2-hexadecenyl]hydroquinone 50, cryst. cellulose 50, cornstarch 30, lactose 17.8, hydroxypropylcellulose 0.2, and Mg stearate 2 mg.

ST isoprenoid deriv prepn antiulcer; phytolhydroquinone deriv prepn
antiulcer; hydroquinone phytol prepn antiulcer

IT Ulcer inhibitors
(isoprenoid derivs.)

IT 50848-64-1P 61977-06-8P 71258-98-5P 77551-14-5P 79577-58-5P
102043-79-8P 123086-31-7P 123086-32-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(prepn. and reaction of, in prepn. of ulcer inhibitors)

IT 119-98-2P 6199-76-4P 10457-66-6P 35175-62-3P 39703-09-8P
71258-97-4P 74166-02-2P 74166-03-3P 76275-68-8P 81557-69-9P
113714-90-2P 119980-00-6P 123086-31-7P 123086-32-8P 123086-33-9P
123086-34-0P 123086-35-1P 123086-36-2P 123086-37-3P 123086-38-4P
123086-39-5P 123086-40-8P 123086-41-9P 123086-42-0P 123086-43-1P
123086-44-2P 123086-45-3P 123086-46-4P 123086-47-5P 123086-48-6P
123086-49-7P 123086-50-0P 123086-51-1P 123086-52-2P 123086-53-3P
123086-54-4P 123086-55-5P 123086-56-6P 123086-57-7P 123086-58-8P
123086-59-9P 123086-60-2P 123086-61-3P 123086-62-4P 123086-63-5P
123086-64-6P 123164-53-4P 123237-20-7P

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(prepn. of, as ulcer inhibitor)

IT 87-66-1, 1,2,3-Benzenetriol 91-16-7, 1,2-Dimethoxybenzene 106-24-1,
Geraniol 119-98-2 120-80-9, 1,2-Benzenediol, reactions 123-31-9,
1,4-Benzenediol, reactions 128-39-2, 2,6-Di-tert-butylphenol 141-97-9,
Ethyl acetoacetate **1113-21-9**, Geranylinalool 4602-84-0,
Farnesol 7541-49-3 13190-97-1, Solanesol 77551-14-5 79577-58-5
123086-37-3 123086-40-8 123086-45-3 123086-47-5 123164-54-5

RL: RCT (Reactant); RACT (Reactant or reagent)

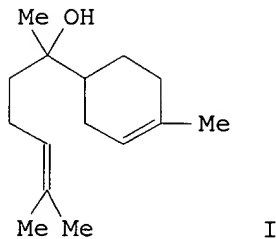
(reaction of, in prepn. of ulcer inhibitors)

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AN 1990:191786 CAPLUS
DN 112:191786
TI Measures of anxiety, retention and **stress** in the rat following treatment with the diterpene sclareol glycol
AU Georgieva, Zh.
CS Inst. Pharmacol. Pharm., Sofia, 1431, Bulg.
SO Methods and Findings in Experimental and Clinical Pharmacology (1990), 12(1), 5-10
CODEN: MFEPDX; ISSN: 0379-0355
DT Journal
LA English
CC 1-11 (Pharmacology)
Section cross-reference(s): 14
AB In a punished drinking test in rats sclareol glycol (SG) decreased the no. of punished responses (proconflict response) while diazepam had the opposite effect; SG antagonized the anticonflict response of diazepam. Post-training administration of SG in rats enhanced retention in active avoidance task evaluated 24 h later. SG produced an increase in plasma ACTH and corticosterone levels in unstressed rats. The **stress**-induced increase in ACTH and corticosterone secretion was potentiated by SG. These data suggest that SG behaves as an anxiogenic, memory-facilitator and perhaps adaptogenic agent. The effects of SG may be mediated by different mechanisms of action (stimulation of adenylyl cyclase or interaction with GABA-ergic and dopaminergic transmitter mechanisms).
ST sclareol glycol anxiety learning **stress**; diterpene anxiety learning **stress**
IT **stress**, biological (ACTH and corticosterone secretion induction by, sclareol glycol enhancement of)
IT Anxiety (from sclareol glycol)
IT Learning (sclareol glycol enhancement of)
IT 38419-75-9, Sclareol glycol
RL: BIOL (Biological study) (anxiety from and learning stimulation by and **stress**-induced increase in ACTH and corticosterone secretion response to)
IT 50-22-6, Corticosterone 9002-60-2, ACTH, biological studies
RL: BIOL (Biological study) (secretion of, sclareol glycol increase of **stress**-induced)

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AN 1979:400383 CAPLUS
DN 91:383
TI Pharmacological experiments with components of chamomile. III.
Experimental animal studies of the ulcer-protective effect of chamomile
AU Szelenyi, I.; Isaac, O.; Thiemer, K.
CS Chemiewerk Homburg, Degussa, Frankfurt/Main, Fed. Rep. Ger.
SO Planta Medica (1979), 35(2), 218-27
CODEN: PLMEA; ISSN: 0032-0943
DT Journal
LA German
CC 1-5 (Pharmacodynamics)
GI



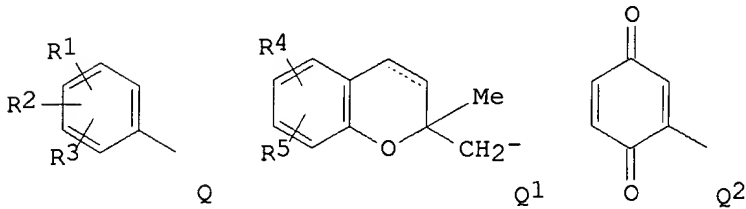
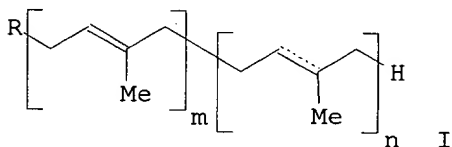
AB (-)-.alpha.-Bisabolol (I) [23089-26-1], a component of chamomile, inhibited the ulcer formation induced by indomethacin, alc., or **stress** in rats, and increased the rate of healing of ulcers caused by HOAc or heat cauterization of the stomach. The chamomile ext. Kamillosan also inhibited the occurrence of alc.-induced ulceration.
ST bisabolol ulcer inhibition; chamomile component ulcer inhibition
IT Ulcer
(bisabolol and chamomile ext. inhibition of)
IT Chamomile
(ext. of, ulcer inhibition by)
IT **23089-26-1**
RL: BIOL (Biological study)
(ulcer inhibition by)

DN 112:36226
 TI Preparation of antiulcer isoprenoid derivatives and pharmaceutical compositions containing them
 IN Yamada, Kazuhiko; Tahara, Yoshiyuki; Toyoda, Masashi; Irino, Osamu; Misaki, Noriyuki
 PA Nisshin Flour Milling Co., Ltd., Japan
 SO Eur. Pat. Appl., 29 pp.
 CODEN: EPXXDW
 DT Patent
 LA English
 IC ICM A61K031-05
 ICS A61K031-085; A61K031-09; A61K031-22; A61K031-12; A61K031-355; C07C050-06; C07D311-72; C07D311-58; C07C039-19
 CC 30-40 (Terpenes and Terpenoids)
 Section cross-reference(s): 1

FAN.CNT 1

| | PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------|---|------|----------|-----------------|----------|
| PI | EP 304842 | A2 | 19890301 | EP 1988-113617 | 19880822 |
| | EP 304842 | A3 | 19910116 | | |
| | EP 304842 | B1 | 19941130 | | |
| | R: BE, CH, DE, ES, FR, GB, IT, LI, NL, SE | | | | |
| | JP 02042030 | A2 | 19900213 | JP 1988-206465 | 19880822 |
| | US 4906669 | A | 19900306 | US 1988-234895 | 19880822 |
| | ES 2067461 | T3 | 19950401 | ES 1988-113617 | 19880822 |
| | KR 9701518 | B1 | 19970211 | KR 1988-10803 | 19880825 |
| PRAI | JP 1987-209214 | A | 19870825 | | |
| | JP 1988-96770 | A | 19880421 | | |
| | JP 1988-206455 | A | 19880822 | | |

GI



AB The title compds. [I; R = Q, Q₁, Q₂; R₁-R₃ = H, OH, alkanoyloxy, alkyl, alkoxy, provided that R₁-R₃ ≠ H, R₄, R₅ = H, OH, alkanoyloxy; m = 0, 1; n = 0-9], useful as antiulcer agents, were prep'd. Hydroquinone in dioxane contg. BF₃.Et₂O was heated 1 h at 40.degree. with 3,7,11,15-tetramethyl-2-hexadecen-1-ol to give, after hydrolysis, 2-(3,7,11,15-tetramethyl-2-hexadecenyl)hydroquinone. At 100 mg/kg orally the latter gave 84.9% inhibition of stress-induced ulcers in male rats. A tablet was formulated contg. 2-[3,7(R),11(R),15-tetramethyl-2-hexadecenyl]hydroquinone 50, cryst. cellulose 50, cornstarch 30, lactose 17.8, hydroxypropylcellulose 0.2, and Mg stearate 2 mg.

ST isoprenoid deriv prep antiulcer; phytolhydroquinone deriv prep
antiulcer; hydroquinone phytol prep antiulcer

IT Ulcer inhibitors
(isoprenoid derivs.)

IT 50848-64-1P 61977-06-8P 71258-98-5P 77551-14-5P 79577-58-5P
102043-79-8P 123086-31-7P 123086-32-8P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)
(prepn. and reaction of, in prepn. of ulcer inhibitors)

IT 119-98-2P 6199-76-4P 10457-66-6P 35175-62-3P 39703-09-8P
71258-97-4P 74166-02-2P 74166-03-3P 76275-68-8P 81557-69-9P
113714-90-2P 119980-00-6P 123086-31-7P 123086-32-8P 123086-33-9P
123086-34-0P 123086-35-1P 123086-36-2P 123086-37-3P 123086-38-4P
123086-39-5P 123086-40-8P 123086-41-9P 123086-42-0P 123086-43-1P
123086-44-2P 123086-45-3P 123086-46-4P 123086-47-5P 123086-48-6P
123086-49-7P 123086-50-0P 123086-51-1P 123086-52-2P 123086-53-3P
123086-54-4P 123086-55-5P 123086-56-6P 123086-57-7P 123086-58-8P
123086-59-9P 123086-60-2P 123086-61-3P 123086-62-4P 123086-63-5P
123086-64-6P 123164-53-4P 123237-20-7P
RL: BAC (Biological activity or effector, except adverse); BSU (Biological
study, unclassified); SPN (Synthetic preparation); BIOL (Biological
study); PREP (Preparation)
(prepn. of, as ulcer inhibitor)

IT 87-66-1, 1,2,3-Benzenetriol 91-16-7, 1,2-Dimethoxybenzene 106-24-1,
Geraniol 119-98-2 120-80-9, 1,2-Benzenediol, reactions 123-31-9,
1,4-Benzenediol, reactions 128-39-2, 2,6-Di-tert-butylphenol 141-97-9,
Ethyl acetoacetate 1113-21-9, Geranylinalool **4602-84-0**,
Farnesol 7541-49-3 **13190-97-1**, Solanesol 77551-14-5
79577-58-5 123086-37-3 123086-40-8 123086-45-3 123086-47-5
123164-54-5
RL: RCT (Reactant); RACT (Reactant or reagent)
(reaction of, in prepn. of ulcer inhibitors)

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